

The Complete Burning of Weapons Grade Plutonium and Highly Enriched Uranium with (Laser Inertial Fusion-Fission Energy) LIFE Engine

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The Complete Burning of Weapons Grade Plutonium and Highly Enriched Uranium with (Laser Inertial Fusion-Fission Energy) LIFE Engine

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Executive Summary

The National Ignition Facility (NIF) project, a laser-based Inertial Confinement Fusion (ICF) experiment designed to achieve thermonuclear fusion ignition and burn in the laboratory, is under construction at the Lawrence Livermore National Laboratory (LLNL) and will be completed in April of 2009. Experiments designed to accomplish the NIF's goal will commence in late FY2010 utilizing laser energies of 1 to 1.3 MJ. Fusion yields of the order of 10 to 20 MJ are expected soon thereafter.

Laser initiated fusion-fission (LIFE) engines have now been designed to produce nuclear power from natural or depleted uranium without isotopic enrichment, and from spent nuclear fuel from light water reactors without chemical separation into weapons-attractive actinide streams. A point-source of high-energy neutrons produced by laser-generated, thermonuclear fusion within a target is used to achieve ultra-deep burn-up of the fertile or fissile fuel in a sub-critical fission blanket. Fertile fuels including depleted uranium (DU), natural uranium (NatU), spent nuclear fuel (SNF), and thorium (Th) can be used. Fissile fuels such as low-enrichment uranium (LEU), excess weapons plutonium (WG-Pu), and excess highly-enriched uranium (HEU) may be used as well. Based upon preliminary analyses, it is believed that LIFE could help meet worldwide electricity needs in a safe and sustainable manner, while drastically shrinking the nation's and world's stockpile of spent nuclear fuel and excess weapons materials. LIFE takes advantage of the significant advances in laser-based inertial confinement fusion that are taking place at the NIF at LLNL where it is expected that thermonuclear ignition will be achieved in the 2010-2011 timeframe.

Starting from as little as 300 to 500 MW of fusion power, a single LIFE engine will be able to generate 2000 to 3000 MWt in steady state for periods of years to decades, depending on the nuclear fuel and engine configuration. Because the fission blanket in a fusion-fission hybrid system is subcritical, a LIFE engine can burn any fertile or fissile nuclear material, including unenriched natural or depleted U and SNF, and can extract a very high percentage of the energy content of its fuel resulting in greatly enhanced energy generation per metric ton of nuclear fuel, as well as nuclear waste forms with vastly reduced concentrations of long-lived actinides. LIFE engines could thus provide the ability to generate vast amounts of electricity while greatly reducing the actinide content of any existing or future nuclear waste and extending the availability of low cost nuclear fuels for several thousand years. LIFE also provides an attractive pathway for burning excess weapons Pu to over 99% FIMA (fission of initial metal atoms) without the need for fabricating or reprocessing mixed oxide fuels (MOX). Because of all of these advantages, LIFE engines offer a pathway toward sustainable and safe nuclear power that significantly mitigates nuclear proliferation concerns and minimizes nuclear waste.

An important aspect of a LIFE engine is the fact that there is no need to extract the fission fuel from the fission blanket before it is burned to the desired final level. Except for fuel inspection and maintenance process times, the nuclear fuel is always within the core of the reactor and no weapons-attractive materials are available outside at any point in time. However, an important consideration when discussing proliferation concerns associated with any nuclear fuel cycle is the ease with which reactor fuel can be converted to weapons usable materials, not just when it is extracted as waste, but at any point in the fuel cycle. Although the nuclear fuel remains in the core of the engine until ultra deep actinide burn up is achieved, soon after start up of the engine, once the system breeds up to full power, several tons of fissile material is present in the fission blanket. However, this fissile material is widely dispersed in millions of fuel pebbles, which can be tagged as individual accountable items, and thus made difficult to divert in large quantities.

This report discusses the application of the LIFE concept to nonproliferation issues, initially looking at the LIFE (Laser Inertial Fusion-Fission Energy) engine as a means of completely burning WG Pu and HEU. By combining a neutron-rich inertial fusion point source with energy-rich fission, the once-through closed fuel-cycle LIFE concept has the following characteristics: it is capable of efficiently burning excess weapons or separated civilian plutonium and highly enriched uranium; the fission blanket is sub-critical at all times ($k_{\rm eff} < 0.95$); because LIFE can operate well beyond the point at which light water reactors (LWRs) need to be refueled due to burn-up of fissile material and the resulting drop in system reactivity, fuel burn-up of 99% or more appears feasible.

The objective of this work is to develop LIFE technology for burning of WG-Pu and HEU. This effort which will enable the development of the LIFE engine for more difficult technical missions, such as the burning DU and LWR SNF, with the simultaneous production of electrical energy or LWR fuel, and will consume a weapons material that poses long-term storage challenges. A point design will evolve for burning WG-Pu and HEU in the LIFE engine without the need for recycling or reprocessing, and with the ability to reach 99% FIMA (fissioned of initial metal atoms).

Tasks and Deliverables

The tasks and deliverables now be executed for NA-22 are summarized in this section, with the budget given in Table 1. During the first phase of this project, emphasis is being placed upon proliferation resistance, engine options, system and performance optimization, fuel design, fabrication and performance, and first wall and structural materials options.

LIFE Materials Tasks

Task 1 – LIFE proliferation resistance

Task 1.1 – Proliferation Resistance Assessment for WG-Pu Fuels

Task 1.1.1 – TRISO Option

Task 1.1.3 – Liquid Fuel Option

Task 1.2 – Proliferation Resistance Assessment for HEU Fuels

Task 1.2.1 – TRISO Option

Task 1.2.3 – Liquid Fuel Option

Task 2 – LIFE engine options and performance optimization

Task 2.1 – Neutronics & Engine Design

Task 2.1.1 – Engine Designs Optimized for Burning WG-Pu Fuels

Task 2.1.1.1 – TRISO Option

Task 2.1.1.3 – Liquid Fuel Option

Task 2.1.2 – Engine Designs Optimized for Burning HEU Fuels

Task 2.1.2.1 – TRISO Option

Task 2.1.2.3 – Liquid Fuel Option

Task 2.2 – Fuel Design, Fabrication & Performance

Task 2.2.1 – Fuel Design & Fabrication Processes for WG-Pu Fuels

Task 2.2.1.1 – TRISO Option

Task 2.2.1.3 – Liquid Fuel Option

Task 2.2.1.5 – Cladding for Abrasion and Corrosion Resistance

Task 2.2.2 – Fuel Design & Fabrication Processes for HEU Fuels

Task 2.2.2.2 – TRISO Option

Task 2.2.2.3 – Liquid Fuel Option

Task 2.2.2.5 – Cladding for Abrasion and Corrosion Resistance

Task 2.3 – First Wall & Structural Materials Options

Task 2.3.1 – First Wall & Structural Materials Options for Solid Fuels

Task 2.3.2 – First Wall & Structural Materials Options for Molten Salt Fuels

Deliverables

Deliverable 1.1 – Report to NA-22 on proliferation implications of the LIFE concept in particular the economic feasibility of >90% burn-up of plutonium

Deliverable 2.1 – Preliminary report to NA-22 on LIFE engine designs for WG-Pu & HEU. Chapters 1, 2, and 4 will be included in the report to NA-22 for completeness, but are being funded elsewhere and not under the current proposal.

Ultimately, reports will include information on Laser Design, Target Design, Engine Design, Target Plant and Injection System, On-Site Solid-State Tritium Handling & Storage, Interim Spent Fuel Storage Containers & Facility; and Preliminary Cost Analysis.

Deliverable 2.2 – Preliminary report to NA-22 on fuel designs, and a quantitative assessment of the performance and lifetime of solid fuel forms. This report will include the following chapters: Documented Materials Selection & Justification; Conceptual Fuel Fabrication Process; Transmutation & Phase Formation; Radiation Effects; Thermomechanical Effects; Corrosion & Environmental Fracture; and Interaction of Solid Fuels with Molten-Salt Coolants.

Deliverable 2.3: A preliminary report will be prepared for NA-22 on fuel designs, and a quantitative assessment of the performance and lifetime of structural and first-wall materials. This report will include the following chapters: Documented Materials Selection & Justification; Conceptual Engine Fabrication & Replacement Process; and Interaction of Structural Materials and Cladding with Molten Salts.

Deliverable 2.4: Preliminary report to NA-22 on the molten salt fuel option, and a quantitative assessment of the performance and lifetime of structural materials and cladding in contact with these fuels. This report will include the following chapters: Available Phase Diagrams for Molten Salt Coolants; Available Phase Diagrams for Molten Salt Fuels; Available Actinide Solubility in Molten Salt Fuels; Preliminary Assessment of Available Fission Product Solubility in Molten Salt Fuels.

Acronyms

ATR Advanced Test Reactor

ABWR Advanced Boiling Water Reactor

ALWR Advanced Light Water Reactor

APWR Advanced Pressurized Water Reactor

BHN Brinell Hardness Number

BWR Boiling Water Reactor

CTE Coefficient of Thermal Expansion

CV Cyclic Voltammetry

EIS Electrochemical Impedance Spectroscopy

FFHR Force Free Helical Reactor

FIMA Fission of Initial Metal Atoms

FLIBE: Molten Fluoride Salt (Li₂BeF₄)

FLINABE Molten Fluoride Salt (LiNaBeF₄)

FLINAK Molten Fluoride Salt (Li-Na-K-F)

HEU Highly Enriched Uranium

HIC Hydrogen Induced Cracking

HFIR High Flux Isotope Reactor

ICF Inertial Confinement Fusion

INL Idaho National Laboratory

IMF Inert Matrix Fuel

IPyC Inner Pyrolytic Carbon (Layer)

LEU Low Enriched Uranium

LIFE Laser Initiated Fusion-Fission Energy

LLNL Lawrence Livermore National Laboratory

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LP Linear Polarization

LWR Light Water Reactor

MD Molecular Dynamic

NIC National Ignition Campaign

NIF National Ignition Facility

NP New Production

NPR New Production Reactor

MHTGR Modular High Temperature Gas-Cooled Reactor

OCP Open Circuit Potential

ODS Oxide Dispersion Strengthened

OPyC Outer Pyrolytic Carbon (Layer)

ORNL Oak Ridge National Laboratory

PWR Pressurized Water Reactor

RN Radionuclide

SCC Stress Corrosion Cracking

SDR Slip Dissolution Repassivation

SHC Solid Hollow Core Fuel

SNF Spent Nuclear Fuel

SSRT Slow Strain Rate Testing

TMT Thermo Mechanical Treatment

TRISO TRI-ISO-tropic Fuel

UTS Ultimate Tensile Strength

WP Waste Package

YMP Yucca Mountain Project

Introduction and Background

Building Upon Anticipated Success of National Ignition Facility

The anticipated success at the National Ignition Facility (NIF) will usher in a new age in fusion physics, with a large number of opportunities for nuclear engineers worldwide to conceive, design and develop fusion, as well as fusion-fission hybrid energy conversion systems. Some concepts have already emerged that promise to provide NNSA with an attractive process for the destruction of special nuclear materials (SNM) such as weapons-grade plutonium (WG Pu) and highly enriched uranium (HEU), while recovering substantial energy to supplement the nation's need for electrical power. Other opportunities exist for the direct burning of natural and depleted uranium, without the need for enrichment. In this case, all possible proliferation scenarios surrounding this promising fuel cycle should be assessed early.

This proposal discusses research opportunities specifically in the area of nonproliferation, initially looking at the LIFE (Laser Inertial Fusion-Fission Energy) engine as a means of essentially completely burning WG Pu and HEU. By combining a neutron-rich inertial fusion point source with energy-rich fission, the once-through closed fuel-cycle LIFE concept has the following characteristics:

- It is capable of efficiently burning excess weapons or separated civilian plutonium and highly enriched uranium;
- The fission blanket is sub-critical at all times ($k_{eff} < 0.95$);
- Because LIFE can operate well beyond the point at which light water reactors (LWRs) need to be refueled due to burn-up of fissile material and the resulting drop in system reactivity, fuel burn-up of 99% or more appears feasible.

Since LIFE is inherently safe, and because neither enrichment nor reprocessing are required, the once-through closed fuel-cycle LIFE approach would significantly mitigate proliferation concerns and provide a sustainable path for worldwide growth of nuclear power for electricity production and hydrogen generation. Moreover, LIFE can efficiently use as fuel excess Pu and HEU and burn them to beyond 99% FIMA (Final Inventory of Metal Atoms). Thus, LIFE provides an interesting path forward for the disposition of these excess weapons-attractive materials. In fact, because of the flexibility and versatility afforded by the combination of fusion and fission, LIFE engines will provide policy makers with options to burn the nation's estimated stockpile of 700 metric tons (MT) of civilian and excess weapons grade Pu (WG-Pu), and highly enriched uranium (HEU). Since LIFE engines can provide approximately 1.1-1.2 GWe-yr/MT of WG-Pu or HEU, this could supply ~ 800 GWe-yrs of electricity. (We note that the US current fleet nuclear power plants provide approximately 110 GWe.) LIFE systems can also burn fertile fuels, and we note that if the nation's stockpiles of SNF and DU were burned in LIFE engines it could supply U.S. electricity needs for more than 1,000 years.

Successful development of LIFE would allow the U.S. to regain global leadership in the development of advanced nuclear energy technologies while mitigating proliferation risks and strengthening American economic competitiveness. Eventually, the work scope should be expanded to assess any and all conceivable nonproliferation opportunities surrounding laser- or potentially heavy-ion-driven fusion-fission hybrid systems.

Objectives for LIFE

There are three primary missions for the LIFE engine, which include the burning of excess weapons-grade plutonium (WG-Pu), highly enriched uranium (HEU), natural and depleted uranium (Natural U and DU), and light-water reactor (LWR) spent nuclear fuel (SNF). The LIFE engine enable the direct burning of such fuels with no isotopic enrichment or chemical separation, with the potential to attain very high burn-up, which is quantified in terms of fission of initial metal atoms (FIMA).

The objective of this work is to develop LIFE technology for burning of WG-Pu and HEU. This effort which will enable the development of the LIFE engine for more difficult technical missions, such as the burning DU and LWR SNF, with the simultaneous production of electrical energy or LWR fuel, and will consume a weapons material that poses long-term storage challenges. A point design will evolve for burning WG-Pu and HEU in the LIFE engine without the need for recycling or reprocessing, and with the ability to reach 99% FIMA (fissioned of initial metal atoms).

Weapons grade plutonium (WG-Pu) and highly-enriched uranium (HEU) can be digested directly into a molten salt, thereby forming a high burn-up liquid fuel for the direct incineration of old weapons materials, as shown in Figure 1. Natural and depleted uranium (Nat-U and DU) can be incorporated into a variety of solid fuels including TRI-structural ISO-tropic fuel (TRISO), baseball or solid hollow core (SHC), inert matrix fuel (IMF), and encapsulated powder fuels, as shown in Figure 2.

Various HEU and WG Pu fueling options will be evaluated. The options that will be evaluated for the HEU and WG Pu mission include: TRISO (tri-isotropic) fuel; or molten salt (Li₂BeF₄ with dissolved PuF₃ or UF₄) fuels. A detailed conceptual process for the fabrication and maintenance of these two fuel forms will be developed, with mass and energy balances, process flow diagrams, equipment sizing, and cost analysis.

Because, in order to explore burning of SNM in LIFE requires an understanding of the overall LIFE system, we describe below the key elements of the LIFE system as well as the proposed scope, deliverables, schedule and cost of the work to be accomplished in the remainder of FY2008.

The final deliverable will be a written report and a briefing package of all aspects of the assessment activities, including an enterprise cost analysis of the economics of burning Pu to

99% FIMA. We will also evaluate and propose experiments to demonstrate the feasibility of extending the life of the critical components of a LIFE engine, specifically first wall and the TRISO-like fuels, to burn SNM and, in particular, plutonium to 99% FIMA. Finally, we will provide a detailed, in depth analysis of the pros and cons of alternative approaches to elimination of excess weapons Pu through burn to greater than 99% FIMA such as accelerator transmutation or magnetic fusion energy based systems.

Application to Burning WG-Pu and HEU

A LIFE plant with a diode-pumped solid-state laser (DPSSL) and NIF-like central hot spot ignition targets is shown in Figure 3 [Moses & Diaz de la Rubia 2008]. In this system, the LIFE engine features a dedicated first wall coolant with a pebble-based multiplier and fuel (Figure 4) [Latkowski, Abbott & Kramer 2008]. After passing through a beryllium neutron multiplier/moderator, the neutrons are coupled to a molten-salt-cooled pebble bed blanket in which the pebbles contain fertile and/or fissile fuel in the form of TRIstructural-ISOtropic (TRISO) particles embedded in a graphite matrix. LIFE is coupled to a helium-based Brayton cycle. A broad range of ICF target-laser combinations exist (Figure 5) [Storm & Lindl 2008]. Different targets would result in different chamber sizes and electric output (Figure 6).

Ultimately, LIFE engines can be developed to the point where they will be able to utilize a wide variety of fuels. Fissile fuels such as low-enrichment uranium (LEU), weapons-grade plutonium (WG-Pu), and highly-enriched uranium (HEU) may be used. Fertile fuels including depleted uranium (DU), natural uranium (NatU), spent nuclear fuel (SNF), and thorium can be used as well.

The burn-up of fission fuel in a LIFE engine can be adjusted as desired with 40 tons of depleted uranium can become nearly 40 tons of fission products (Table 2). When burning DU, several isotopes will be fissioning in the LIFE engine (Figure 7). Since the fission of DU generates a fuel that is predominately plutonium after a few years, the characteristics of DU and WG-Pu burners are very similar at long times. Knowledge of machines designed to burn plutonium is pertinent to the design of systems for burning DU, and vice versa. LIFE fuel burn-up can be adjusted as desired (Figure 8) [Latkowski, Abbott & Kramer 2008].

The TRISO-based LIFE fuel can achieve a total actinide burn-up of more than 99 percent without refueling, and has an energy release of about 1 MW-day per gram fissioned. In an optimized LIFE engine, this translates into approximately 1.1 to 1.2 GWe-year per ton of fuel burned. The greater than 99% burn-up also means that each ton of fuel burned in LIFE would contain less than 10 kg of actinides. This compares quite favorably to the 0.97 tons of actinides that remain for each ton of fuel burnt in a typical LWR.

This has major consequences for the waste disposal required for LIFE power plants. Once the system reaches full power, several tons of fissile material is present in the blanket, but this is widely dispersed in millions of fuel pebbles. Any collection of pebbles that is capable of

providing a significant quantity (SQ) of fissile material (defined as 8 kg for ²³⁹Pu) is self-protecting in that it provides a dose rate of at least 1 Gy/hour (100 rad/hour) at a distance of 1 meter. At the end of the burn cycle, the full collection of pebbles contains less than one SQ of ²³⁹Pu. Given the large fraction of fission products and low fraction of long-lived actinides in LIFE spent fuel, it may be possible to dispose of the spent fuel in a simpler manner than expected for LWR waste.

LIFE can burn excess plutonium in a high-gain blanket, and is being explored for NA-22 (Table 3, Figure 9). This conceptual system is assumed to be fueled with 7 tons of weapons-grade plutonium (WG-Pu). It is further assumed that: TRISO kernels are blended 80% ZrC + 20% Pu; TRISOs are loaded with 400 wppm boron as burnable poison; fusion power is 375 MW (37.5 MJ at 10 Hz); the flat-top thermal power is 3000 MW; and the blanket gain is \sim 8. When fueled with weapons-grade Pu or HEU, LIFE will typically be fueled with \sim 6 or 7 metric tons and with \sim 350-400 MW fusion provide 3000 MWth or 1350 MWe for about 6 years while burning the Pu to over 99% FIMA.

An early LIFE demonstration unit, burning WG-Pu, will enable the development of an advanced power system capable of helping provide much of the Nation's future energy needs, with substantial reductions in the generation of high-level radioactive waste (Figure 10). Specific system improvements that are anticipated include the advent of fast-ignition ICF, advanced solid and molten salt fission blankets, and advanced structural materials capable of operating at higher temperatures. The WG-Pu mission has the potential of enabling the hybrid LIFE systems, and will also enable pure fusion, since LIFE is a practical and realizable precursor for pure fusion (Figure 11).

In summary, we have discussed a concept for a hybrid fusion-fission energy engine. Significant challenges in demonstrating the physics of IDFI, development of advanced materials and DPSSL technologies remain. However, recent progress on NIF at LLNL and on the physics and technologies of inertial confinement fusion, DPSSL technologies and materials science for applications in the nuclear environment in the US and elsewhere around the world promise to bring us closer to realizing the longstanding vision of fusion-fission hybrids. Starting with a laser fusion driver with gain of ~ 50 , such a device would be capable of burning large quantities of spent nuclear fuel, natural or depleted uranium or weapons-grade Pu to generate GW of thermal power in a subcritical fission blanket. It would therefore provide pathways to energy generation in a closed nuclear fuel cycle and to eliminating the excess inventory of spent nuclear fuel on the way to pure fusion energy.

Other Non-Proliferation Issues Associated with Dual-Use Technology

Other opportunities exist for the direct burning of natural and depleted uranium, without the need for enrichment. In this case, all possible nonproliferation scenarios surrounding this promising fuel cycle should be assessed early.

The concept of a Laser Inertial Fusion-Fission Energy (LIFE) engine has the potential to generate GWs of thermal power while closing the nuclear fuel cycle and provides significant safety and nonproliferation advantages over current and proposed nuclear energy options. By combining a neutron-rich inertial fusion source with energy-rich fission, the once-through closed fuel-cycle LIFE concept has the following characteristics:

- It eliminates the need for enriched nuclear fuel and can operate on depleted uranium or thorium;
- Because LIFE can operate well beyond the point at which light water reactors (LWRs) need to be refueled due to burn-up of fissile material and the resulting drop in system reactivity, fuel burn-up of 99% or more appears feasible, and LIFE thus eliminates the need for refueling and reprocessing facilities;
- It is capable of burning Spent Nuclear Fuel (SNF) to over 99% burn without chemical separation of actinide streams;
- It minimizes future requirements for deep underground geological waste repositories and minimizes actinide content in the end of life nuclear waste below DOE's attractiveness level E (the lowest).

LIFE systems can also burn fertile fuels, and we note that if the nation's stockpiles of SNF and DU were burned in LIFE engines it could supply U.S. electricity needs for more than 1,000 years. Since LIFE is inherently safe, and because neither enrichment nor reprocessing are required, the once-through closed fuel-cycle LIFE approach would significantly mitigate proliferation concerns and provide a sustainable path for worldwide growth of nuclear power for electricity production and hydrogen generation. LIFE offers several technical and nonproliferation advantages over other systems.

- 1) The LIFE engine is deeply subcritical at all times and in all configurations.
- 2) It is able to passively accommodate either loss of flow or total loss of coolant accidents without a significant temperature rise in the fuel pebbles.
- 3) LIFE eliminates the need for costly uranium enrichment and refueling, resulting in sizeable cost savings as well as significantly mitigating nuclear proliferation concerns. A nation operating LIFE engines will never have the need to build nuclear enrichment or reprocessing facilities.
- 4) The LIFE engine extracts virtually 100 percent of the energy content of its fuel (versus approximately 0.5 percent of the energy in the ore required to make fuel for a typical LWR.
- 5) LIFE drastically minimizes requirements for geologic waste repositories, saving hundreds of billions of dollars in construction and maintenance costs. LIFE offers a way to "burn to a nuclear crisp" all the SNF now destined for transportation to and storage in Yucca Mountain as well as the huge supply of depleted uranium that exists now and will be created in the decades ahead.
- 6) In addition to burning natural uranium, a LIFE engine can use for its fuel the two waste streams associated with the present and future fleet LWRs, Spent Nuclear Fuel (SNF) and depleted uranium left over from the process used to enrich uranium.
- 7) When compared with existing and other proposed future nuclear reactor designs, the LIFE engine exceeds alternatives in the most important measures of proliferation resistance. By

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integrating fuel generation, energy production, and waste minimization into a single device, the LIFE engine is intrinsically highly proliferation-resistant. The engine needs no refueling during its lifetime. It requires no removal of fuel or fissile material generated in the reactor. It leaves no weapons-attractive material at the end of life.

We anticipate that these nonproliferation and technical advantages will result in cost-competitive operation and generation of electricity and process heat for hydrogen generation. Coupling the cost avoidance associated with eliminating uranium enrichment, fuel reprocessing and minimizing requirements for future geological waste repositories with anticipated advances in diode-pumped solid state laser and target technologies (in particular the continued reduction in the cost of high power diodes and the use of high-volume manufacturing technologies for the targets), our initial projections for capital and operating costs show that the cost of electricity from the LIFE engine should be competitive with the projected cost of electricity for advanced nuclear reactor options envisaged for the 2030 time frame.

Key Challenges for Development of LIFE

The key technical challenges associated with the development of a hybrid fusion-fission energy engine can be divided into those related to the fusion source (obtaining adequate target gain and the high-repetition-rate laser) and those associated with the operation of the fission blanket. The approaches to resolving near term issues associated with these technical challenges are given below. This proposal is concentrating on the last two challenges: the first-wall fission blanket and the issues associated with deep-burn fuel. In reviewing the key technical challenges, it is important to note:

- 1) Laser-driven fusion ignition will be attempted on NIF starting in 2010. This is a sine qua non condition for success of LIFE.
- 2) Development of Fast Ignition and planning for experiments to demonstrate the validity of the physics are underway in the U.S., Japan, the EU, and China.
- 3) NIF will execute one laser shot every few hours. A hybrid engine needs to execute 10 shots per second. This requires solid-state lasers using low-cost diodes as the enabling technology. The expectation is that within the next decade diode costs will decrease significantly, driven by the consumer products market.
- 4) Requirements for NIF operation will be about 700 to 1,000 targets per year. The hybrid engine will require 100 million, low-cost targets per year. Technologies for low-cost, large-volume manufacturing can be adopted from other industries. Target injection and laser pointing techniques are under development at General Atomics and Lawrence Livermore National Laboratory.
- 5) The first wall of the fission blanket will be exposed to large, damaging fluxes of fast neutrons, ions, and x rays. Ongoing development in Japan, the EU and the U.S. of new structural steels for fusion and fission reactors provides new materials that are suitable to the hybrid engine application. Ultimately, experimental work will be done, looking at the effects of radiation and thermal pulsing on both the tungsten material, and the interface between the tungsten armor and the ODS or TMT steel structure. Such thermal pulsing can be simulated in the laboratory with (1) laser flash heating of planar surfaces and (2) ohmic heating of samples with current pulses.
- 6) The fission fuel must be capable of withstanding the extreme environment of the fission blanket for decades. This requires advanced materials capable of performing safely at the elevated temperatures and radiation fluxes present in the engine. There is currently much development worldwide in advanced fuel forms capable of burning more than 90% of their initial nuclear fuel without having to recycle or reprocess. Detailed finite element models of TRISO fuels have been developed with LLNL's ALE3D code, and have been successfully benchmarked with published IAEA standards for TRISO simulation. Thus far, sufficient detail has been incorporated into these models to account for SiC capsule wall stress due to fission gas accumulation inside the capsule, thermal pulsing due to laser firing, and debonding between layers. Work has also been done on thermochemical modeling of fission product interaction with the SiC capsule.

LIFE Development Pathway

Because of the ability to separate functional subsystems of the LIFE engine, the science and technology for an integrated demonstration of the LIFE engine can be performed at the modular level in appropriately-scaled facilities. For example, the physics of the indirect fusion fastignition target and demonstration of the required fusion gain can and would be done independently of the LIFE engine and the fission process. In fact, we expect to demonstrate the physics of fast ignition on NIF. Similarly, target production and target delivery can be developed and demonstrated independently, and at the modular level. The DPSSL technology will be demonstrated at the 100 kW beam line level.

We believe that a proof-of-principle LIFE fission cycle could be demonstrated in the next decade if properly funded and a proof-of-principle LIFE fission cycle (breeding up from depleted U through high burn-up) could be demonstrated with 20% scale LIFE engine using depleted U in less than three years after that.

The objective of this proposal is to develop a preliminary assessment of the feasibility of LIFE engines designed to burn plutonium and highly enriched uranium. That is, the LIFE engine driven by 300-400 MW of fusion power to generate ~3000 MW of thermal power from WG-Pu and HEU fuels. This proposal will concentrate on three specific areas that are critical to prove the applicability of the LIFE concept to nonproliferation. These three areas are:

- A systems evaluation of the LIFE concept including the economic feasibility of complete burn-up (>90%). This will also include a survey of other fission-fusion concepts, comparing and contrasting LIFE's risks and benefits with regards to other concepts.
- The technical issues associated with the survival of components in the harsh environment of the fusion-fission process. In particular, mitigation of early removal of the TRISO fuel before complete burn-up.
- Design of the fission blanket to maximize the benefit of LIFE for nonproliferation purposes.
- Below we describe the key elements of the LIFE system addressed in this proposal as well as
 the proposed scope, deliverables, schedule and cost of the work to be accomplished in the
 remainder of FY2008.

Task 2 - LIFE Engine Options, Materials and Performance

Task 2.1 - Systems Evaluations

Systems modeling and analyses are required to evaluate various design, performance and global impacts of LIFE with specific emphasis on assessing the proliferation resistance of LIFE for burning WG-Pu. Dynamic enterprise models have been developed by LLNL to examine features of the U.S. nuclear infrastructure life-cycles including themes related to the Nuclear Weapons Complex Transformation as well as the Global Nuclear Energy Partnership (GNEP) Program. Nuclear energy is only one part of a larger system of energy supply and this set of models will be adapted to include LIFE plants as they penetrate into the energy system in the context of various future US and global energy growth scenarios.

Associated carbon emissions reductions and potential impacts on global climate change will also be included. The energy system model will include: sources, conversions and services driven by demand-based projections from the Energy Information Agency (EIA) and the International Energy Agency (IEA). Carbon emissions reductions will be referenced to the Inter-Governmental Panel on Climate Change (IPCC) data.

The enterprise model will be capable of tracking material flow (e.g., fuel input, spent fuel, waste, etc.) and infrastructure needs (e.g., fuel factories, power plants, reprocessing plants, long-term storage facilities, etc.) over time. Underlying the top-level (communications-level) enterprise model are more detailed cost and performance systems models for the elements contributing to the nuclear energy mix (e.g., fission plants, LIFE plants, enrichments plants, reprocessing facilities, etc.). A detailed model for LIFE, including the fuel fabrication facility, laser, target factory, engine, power conversion system and balance of plant, will be developed and incorporated in the enterprise model.

Various systems design studies will be completed at both the power unit systems model level (i.e., a single LIFE plant) and at the broader energy system market level for various scenarios of technology introduction time and growth rate. These studies will help identify key design trade-offs and areas of R&D with high leverage for design improvement within the context of pre-LIFE introduction carbon and energy system improvements. Finally, these studies will quantify the potential benefits of the LIFE technology in terms of reduced proliferation risk, reduced long-lived nuclear waste and the impact of LIFE on local, US and global carbon emissions reductions with potential climate change.

The systems studies will include a semi-quantitative comparison to other advanced nuclear energy technologies, such as the GNEP/AFCI fuel cycle and reactor technologies, Gen-IV reactors, and pure fusion. Topics for comparison include: cost, resource utilization, waste minimization and proliferation resistance. System studies will also explore synergistic R&D opportunities with these other technologies, where common issues permit shared research.

Performance modeling falls into the several broad categories, including but not limited to nuclear physics and radiation transport, mechanical design of the engine, including thermal hydraulics, modeling and analysis of the first wall and structure, accounting for their mechanical behavior, as well as various modes of materials degradation.

The performance characteristics of various types of LIFE engines, capable of burning both depleted uranium and plutonium, are summarized Table 4. Plants based upon "hot-spot" ignition require only a compression laser, while plants based on "fast ignition" require both compression and ignition lasers.

Nuclear Reactions

The core of the LIFE Engine is an ICF-driven fusion system that generates high numbers and fluxes of 14 MeV neutrons. These neutrons transport into a blanket containing both fuel pebbles and tritium breeding material. In some cases, a secondary chamber may be utilized to produce tritium that can be fed into targets for the primary chamber. Our current capability to complete burn calculations suffers from a lack of spatial resolution and temperature-dependent data. Partial coupling of neutronics and heat transfer calculations will be implemented during FY2008. The LIFE code suite will be updated to enable nuclear burn to occur in multiple regions and materials within a given model. Additional physics will be added, if necessary. Also, we will continue to improve overall performance in order to increase productivity. Nuclear data libraries will be upgraded for both the radiation transport and activation codes. Using the updated capabilities and data libraries, we will update and improve the LIFE Engine designs for both power production and materials testing options. Benchmarking evaluations and validation experiments will be designed.

The neutrons for driving the sub-critical fission blanket are produced by implosion of the DT capsule in the inertial-confinement fusion (ICF) target in the center of the engine. The well-known D-T reaction in the target consumes tritium and deuterium:

$${}_{1}^{3}T + {}_{1}^{2}D \longrightarrow {}_{2}^{4}He + {}_{0}^{1}n + 17.6MeV$$

The energy imparted to the neutron in the D-T reaction is approximately 14 MeV. This energy is sufficient to make the neutron relativisitic [Equation 4.13, Foster & Wright 1973].

When the neutrons from the fusion target enter the coolant (or liquid fuel), they react with ⁶Li to produce ³T which can be separated, stored, and eventually fed back to the LIFE engine. The lithium transmutation reaction is [Equation 4.14, Foster & Wright 1973]:

$$_{3}^{6}Li+_{0}^{1}n\longrightarrow _{3}^{7}Li\longrightarrow _{1}^{3}T+_{2}^{4}He$$

Natural lithium consists of 92.58% ⁷Li and 7.42% ⁶Li. According to the published literature, natural lithium has an absorption cross-section for thermal neutrons of approximately 70b, while

 7 Li has a neutron absorption cross-section of approximately 0.033b. The transmutation of 6 Li $^{+}$ in the FLIBE produces 3 T $^{+}$, which is very corrosive.

Approximately 99.3% of the world's natural uranium is ²³⁸U, which cannot be fissioned by thermal neutrons, but can be converted to fissile material with the fast neutrons produced in fusion-fission hybrids such as the LIFE engine. Fast (14 MeV) neutrons from the fusion targets will be used for the conversion of fertile nuclei (natural and depleted uranium, and spent lightwater reactor fuel) via the following reactions [Equations 4.10a and 4.10b, Foster & Wright 1973]:

$$^{238}_{92}U + ^{1}_{0}n \longrightarrow ^{239}_{92}U \longrightarrow ^{239}_{93}Np + ^{0}_{-1}e$$

$$^{239}_{93}Np \longrightarrow ^{239}_{94}Pu + ^{0}_{-1}e$$

Neutron capture produces ²³⁹U, which decays to ²³⁹Np by an initial beta emission, which in turn decays to ²³⁹Pu by a second beta emission. Thus, the LIFE engine burns plutonium during most of its service period. In a similar manner, such hybrids can be used to support a thorium-based fuel cycle. Neutron capture converts ²³²Th to ²³³Th, which decays to ²³³Pr by an initial beta emission, which in turn decays to ²³³U by a second beta emission.

It is possible to segment the fission blanket such that the innermost subset of pebbles burns at an accelerated rate while the outer pebbles provide most of the thermal power. Once the innermost pebbles are fully burned, each successive layer would be promoted forward, and fresh fuel would be loaded at the back of the blanket. By operating in this mode, it is possible to extend the flat portion of the power curve for as long as is desired. A key question that must be answered is if the segmentation will naturally occur without the addition of structural material (e.g., flow channels). Such structural material is undesirable in that it would attenuate and absorb neutrons. This question will be addressed in a computational setting and a confirming experiment will be designed. Nuclear burn calculations will be completed for a variety of segmented blanket designs.

Design, Modeling and Analysis

We will use the RadHeat code to transport x-rays and ions through a neutral background gas to see what reaches the first wall as a function of time. Separate calculations will estimate the level of secondary x-ray generation resulting from reverse Bremsstrahlung during ion stopping. These secondary x-rays will be added to the baseline x-ray output to determine the integrated effect upon the first wall. The above calculations will be updated as new target output results are made available by the LLNL target design team.

The pulsed x-rays will cause LIFE's tungsten-armored first wall to experience thermal transients. It is critical to the survivability of the first wall that we understand the implications of these temperature spikes, including the induced stress, strain, cracking and bond fatigue issues. An

experiment will be designed that will employ a rep-rated driver to reproduce relevant time-temperature profiles in the candidate first wall materials at an accelerated rate. If such experiments can be carried out with sufficient fidelity using a multi-kHz laser, then it would be possible to conduct complete lifetime testing in very short periods.

LIFE Engine Materials & Fuels

The fusion-fission chamber is the heart of the LIFE engine, and is shown in Figure 12. Materials of construction fall into several broad categories, including: (1) lasers and optics, (2) fusion targets, (3) tungsten first wall, (4) neutron multiplication blanket, (5) sub-critical fission blanket, (6) structural and cladding materials, (7) coolants and/or liquid fuels, (8) reflector, and (9) balance of plant. Issues related to lasers and optics, as well as fusion targets are discussed elsewhere. This collection of topical reports focuses on fission fuels, structural and first wall materials, the neutron multiplier, coolants and some special materials such as metal tritides required for the balance of plant. This report focuses on the interaction of (6) structural and cladding materials with (7) coolants and/or liquid fuels. As shown in Figure 13, the operating windows for these materials are bounded by radiation embrittlement at lower temperatures, and by thermal creep at higher temperatures. The operating conditions for LIFE have been selected to enable the survivability of known materials of construction.

We will deliver an early predictive model, a detailed plan for experiments necessary to calibrate and validate model parameters and initial experiments to enable predictions and credible materials selections. Specific issues that must be dealt with include: stresses associated with the accumulation of fission gases within the particles of fuel surrounding the fusion target area; pulsed heating of the fuel and first wall, and the associated mechanical stresses; high-temperature creep of structural materials; corrosion and environmental cracking of those reactor components exposed to the high-temperature molten-salt coolant and tritium breeding blanket; radiation damage on fuels, structural and first wall materials; the creation of appropriate test environments for materials within the laboratory, including extreme neutron fluxes with representative energy spectra, rapid thermal pulsing with homogeneous heating; exposures to high-temperature lithium-beryllium-fluoride molten salts, and special problems associated with the use of beryllium multipliers in such applications. Planning, modeling and early experimental work will be used to begin solving materials problems currently believed to challenge the LIFE reactor concept. A detailed research and development strategy will be formulated, thereby enabling the team to attack crucial materials problems in an efficient and comprehensive manner.

The initial fabrication of prototypical fuels by an experienced industrial partner will be initiated, incorporating those design changes necessary to allow the fuels to survive higher burn-up and neutron dose. The proposed developmental work will leverage the extensive knowledge base within the Department of Energy that is now being used in developing the Advanced Gas Reactor (AGR), to provide the technical basis for creating fuels capable of even higher burn-up, and capable of accommodating a much broader range of uranium feeds. This section of the

proposal describes what will be done to address materials challenges, options and performance optimization.

Tungsten First Wall

The high-energy (14 MeV) neutrons generated within the deuterium-tritium (DT) filled fusion targets will travel outward, first encountering a layer of solid tungsten armor supported by a low-activation ODS ferritic steel substrate. This "first wall" should be able to withstand bombardment by the hard-spectrum neutrons emitted by the target, as well as the high temperatures resulting from the absorption of X-rays that accompany the neutrons. The gas that fills the chamber is expected to offer protection from ions.

These materials will be subjected to a severe radiation and high-temperature environment, as shown in Figure 14. Energy from ICF target carried by: neutrons (14 MeV); Ions (< 100 keV); X-rays (< 100 keV); gamma ray contribution small. The target chamber and beam path will be filled with xenon gas at an atomic density of approximately $3 \times 10^{16} \text{ cm}^{-3}$. This gas absorbs a significant portion of the x-ray energy and stops all ions emitted from the indirect-drive target. The hot gas cools via radiation on a timescale sufficiently long to prevent damage to the tungsten first wall.

Specific requirements for the tungsten first wall are summarized in Table 5. This material will have to withstand temperatures of 750 to 1300° C, and bombardment with 8.00×10^{22} neutrons per square centimeter per year (total of 3.20×10^{24} n/cm²), with 2.89×10^{22} neutrons per square centimeter per year (total of 1.16×10^{24} n/cm²) being fast neutrons. The corresponding damage will be approximately 10 displacements per atom per year (total of 400 dpa), with helium and hydrogen at 10 and 22 atomic parts per million per year (total of 400 and 880 appm), respectively.

Neutron Multiplication Blanket

After passing through the tungsten first wall, the neutrons from the target will then enter a layer of beryllium which serves as a neutron multiplier. Be(n,2n)Be reactions moderate their energy and generate approximately 2 neutrons for every one absorbed. The moderated and multiplied neutrons will then strike the next layer, a sub-critical fission blanket. The baseline fission fuel will be in the form of approximately 2-cm diameter pebbles containing a large number of smaller TRISO particles embedded in a graphite inert matrix. Other fuels are also being investigated with the objective of ultra-deep burn (99.9% FIMA).

Specific requirements for the beryllium neutron multiplier are summarized in Table 5. This material will have to withstand temperatures of 650 to 750° C, and bombardment with 7.87×10^{22} neutrons per square centimeter per year (total of 3.15×10^{24} n/cm²), with 2.28×10^{22} neutrons per square centimeter per year (total of 9.12×10^{23} n/cm²) being fast neutrons. The corresponding damage will be approximately 13 displacements per atom per year (total of 520 dpa), with

helium and hydrogen at 5,000 and 51 atomic parts per million per year (total of 200,000 and 2,040 appm), respectively.

Current fusion reactor concepts consider the application of beryllium as neutron moderator and neutron multiplier [Ursu 1985, Section 12.5.3, p. 434]. The 14-MeV neutrons from the ICF target bombard beryllium in the neutron multiplication blanket where Be(n,2n)Be reactions moderate their energy and generate approximately 2 neutrons for every one absorbed. Beryllium is one of the best solid moderators and is used either as metallic beryllium or as beryllium oxide. Beryllium has a low absorption cross section, a high scattering cross section, and a high melting point of 885°C (1158K). Beryllium is a light metal with an elastic modulus slightly superior to that of steel, and does not readily oxidize in air.

With the advent of fusion and fusion-fission reactors in the future, beryllium is expected to become and increasingly important material. Published estimates indicate that a fusion reactor generation capacity of one-million megawatts will require approximately 4.6×10^7 kilograms of beryllium. The availability of this strategic material is discussed by Shedd [2007].

Beryllium is stable under the modest neutron irradiations that have been encountered in conventional nuclear systems, but may be problematic at the fluencies expected in the LIFE engine [Ursu 1985]. Its exposure to neutron fluencies up to 2.3×10^{21} n/cm² does not entail any sizeable dimensional change, nor does its electrical resistivity and corrosion resistance changes. As for the other properties, the thermal conductivity decreases slightly, the tensile strength increases from 500 MPa to about 714 MPa, whereas the elongation decreases from 1.4 to 0.3 percent. Notably, the hardness increases by about 30%. Also, the generation of helium as a result of irradiation causes beryllium to become brittle.

Beryllium and its salts, including FLIBE and FLINABE are considered very toxic and will have to be handled very carefully. Assuming an eight-hour work day, the maximum allowable concentration of beryllium aerosol in a fusion reactor facility should be no more than 2 micrograms per cubic meter. Exposure of he public should be no more than 0.01 micrograms per cubic meter.

Fission Blanket

An enhanced TRISO fuel particle with a more robust silicon carbide capsule than conventional fuel particles enables fission-gas containment at very high burn-up. This TRISO configuration is being considered as one possible high burn-up fuel option for the LIFE engine. The enhanced TRISO will have to endure more radiation damage than conventional TRISO fuels. The experience base for TRISO fuels is limited, with 8 to 20% FIMA demonstrated with low enriched uranium (LEU), and 79% FIMA demonstrated with highly enriched uranium (HEU) [Petti et al. 2004]. These LEU and HEU fuels experienced only approximately 3.5×10^{21} n/cm², but at a relatively high temperature of 1100 to 1200°C. In contrast, LIFE fuels will experience a much higher neutron dose of approximately 1.2×10^{23} n/cm², but at a much lower temperature of

about 700°C. The temperature spike in the fuel pellets that result from the pulse of neutrons entering the fission blanket every 1/20th of a second is approximately 20°C.

Specific requirements for the silicon carbide capsule and graphitic material in the TRISO fuel are summarized in Table 5. The silicon carbide capsule will have to withstand temperatures of 750 to 800° C, and bombardment with 2.97×10^{21} neutrons per square centimeter per year (total of 1.18×10^{23} n/cm²), with 7.66×10^{20} neutrons per square centimeter per year (total of 3.06×10^{22} n/cm²) being fast neutrons. The corresponding damage will be approximately 6 displacements per atom per year (total of 240 dpa), with helium and hydrogen at 88 and 139 atomic parts per million per year (total of 3,520 and 5,560 appm), respectively. The graphitic materials will have to withstand temperatures of 750 to 800° C, and bombardment with 2.97×10^{21} neutrons per square centimeter per year (total of 3.06×10^{22} n/cm²), with 7.66×10^{20} neutrons per square centimeter per year (total of 3.06×10^{22} n/cm²) being fast neutrons. The corresponding damage will be approximately 3 displacements per atom per year (total of 120 dpa).

Fertile & Fissile Material Options

Several types of fertile and fissile materials were considered during the conceptual design of the LIFE engine. Solid uranium fuels could be fabricated from nitrides, oxides, carbides, oxycarbides, silicides, aluminides and metallic alloys. The density and melting points of candidate fissionable fuels are summarized in Table 6, and are listed from the highest to the lowest melting point. Oxides, nitrides and carbides have the highest melting points. Commercial TRISO fuels use a mixed ceramic phase, uranium oxy-carbide (UOC), which appear to minimize thermal migration effects of the kernel.

Compounds such as UF₄ and Th₄ are soluble in molten salts comprised of lithium fluoride and beryllium fluoride, such as FLIBE [Ursu, Chapter 7, p. 247, 1985]. These dissolved uranium and thorium salts can therefore be used as homogenous liquid-phase fuels, which are immune to radiation damage, but are prone to the precipitation of metallic fission products and plutonium (PuF₃). Continuous on-line reprocessing will be required for such fuels, and the potential risk of proliferation may be higher than that of a solid fuel. An obvious exception would be the burning of weapons grade plutonium, where the threat of proliferation will only diminish. If these materials become oxidized, problematic volatile compounds can form. The gaseous UF₆ species is very well known, and could form during the inadvertent oxidation of UF₄. Such volatilization could be controlled through electrochemical control and manipulation of the uranium oxidation state.

Concepts for blanket/coolant materials have evolved over several decades. That evolution is discussed in the book by Ursu [1985], as well as in earlier articles by Grimes and Cantor [1972] and others. The early evolution of these concepts is briefly summarized as follows:

• 1957 (UKAEA, Harwell): $^{238}U - ^{6}Li$

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- 1957 (KAPL): UO₂SO₄
- 1965 (MIT): LiF-BeF₂-UF₄ and LiF-UF₄
- 1969 (MIT): 71%LiF-2%BeF₂-27%ThF₄
- 1970 (MIT): ²³⁵U-²³³U-²³⁹Pu-⁶Li
- 1972 (PNL): Natural Uranium
- 1972 (Grimes et al. at ORNL): Li₂BeF₄ and LiNaBeF₄

Molten Salt Coolant

Clad fuel pebbles formed from several thousand TRISO fuel particles, or an alternative solid fuel such as SHC or IMF are immersed in a molten fluoride salts that carries away heat to drive electrical generators. The very high volumetric heat capacities of these salts allow the fission blanket to be compact and have high power density when coupled to a point source of fusion neutrons (fusion target) [Sviatoslavsky et al. 2004]. Preferred coolants are FLIBE (Li₂BeF₄) in the primary coolant loop and FLINABE (LiNaBeF₄) in the secondary coolant loop. Note that FLIBE is a binary mixture of lithium and beryllium fluorides (2LiF + BeF₂ = Li₂BeF₄). The FLIBE input temperature is 620°C and the exit temperature for this design is 680°C. The transmutation of lithium in these coolants produces tritium for the steady stream of fusion targets that must be fed to the LIFE engine, thereby making the system self sufficient in tritium. Unfortunately, this reaction also produces very corrosive hydrofluoric acid species (HF and TF), which can rapidly degrade structural and cladding materials. In the case of homogenous liquid fuels, UF₄ can be dissolved in these molten salt mixtures.

High-temperature liquid-phase coolants appropriate for heat transfer in nuclear power systems such as the LIFE engine fall into two broad categories, liquid metals and molten salts. Liquid metal include Hg, Na, K, Pb, Bi, Pb-Bi, Sn, Li, Li-Al, Pb-Li, Sn-Li, and others. Molten salts include FLIBE ($2\text{LiF} + \text{BeF}_2 = \text{Li}_2\text{BeF}_4$), FLINABE ($\text{LiF} + \text{NaF} + \text{BeF}_2 = \text{LiNaBeF}_4$), FLINAK (LiF + NaF + KF) and others (for example, $^{24}\text{NaBF}_4 + ^{24}\text{NaF}$). FLIBE has been proposed for use in the primary coolant loop of the LIFE engine, while beryllium-free FLINAK has been proposed for use in the secondary loop. The lithium in the primary loop coolant will enable tritium breeding, while the beryllium helps moderate neutrons.

Molten salt coolants are less prone to catch fire than the liquid metal coolants, and can be used to breed tritium via the transmutation of lithium. A binary mixture of BeF₂ and LiF produces the eutectic composition Li₂BeF₄, which has a relatively low melting point of less than 400°C, as shown in Figure 34. The formula weight, density, melting point and boiling point of several of these candidates are summarized in Table 8 [Williams et al. 2006; Moir et al. 1985; Wright and Foster 1973; Rosenthal et al. 1971]. Pure FLIBE has relatively low viscosity, comparable to kerosene at ambient temperature, and therefore would be expected to flow well.

Note that several other programs have investigated the use of molten fluoride salts. FLIBE is considered as a candidate material for tritium breeding in a fusion liquid due to its chemical stability and low electrical conductivity. The *Jupiter-II Irradiation Tests for Fusion Research* also focused on the use of FLIBE [Petti et al. 2006]. It was selected for the conceptual design of the *Force Free Helical Reactor (FFHR)*. The tritium release behavior and the corrosion of the structural materials by FLIBE were key to success in this application. In this particular design, a ferritic steel (Fe-9Cr-2W) and a vanadium alloy (V-4Cr-4Ti) are the candidate materials for the blanket structure [Nishimura et al. 2001].

Investigators at the University of Tokyo (YAYOI) placed a Monel crucible with 100 to 300 grams of FLIBE in a fast-neutron reactor and heated the salt to 873K [Nishimura et al. 2001]. The neutron flux during these irradiations was 10⁸ to 10⁹ n cm⁻² sec. At hydrogen partial pressures above 1000 Pa, the primary tritium-bearing species responsible for tritium release is gaseous diatomic HT. These investigators determined the normalized release rates for tritium in flowing stream of helium purge gas with various concentrations of hydrogen. The normalized tritium release rate dropped dramatically below hydrogen partial pressures of 1000 Pa. Steady state was achieved after approximately 120 to 140 minutes of purging.

Unfortunately, the transmutation of lithium in this coolant produces very corrosive tritium fluoride species (TF), which behaves like hydrofluoric acid and can rapidly degrade structural materials. Thermal neutrons react with Li⁺ and convert it to T⁺, which then reacts with fluoride anions to form gaseous TF. Exchange reactions between tritons (T⁺) and diatomic hydrogen (H₂) formed protons (H⁺) and diatomic HT. The corrosion of structural components of the LIFE engine in contact with this coolant must be understood.

It may be possible to control the release of tritium from the FLIBE blanket through the manipulation of electrochemical oxidation and reduction reactions, using an on-line electrolytic cell. The reduction of tritons (T^+) in such a cell would simultaneously remove the threat of acidic TF, while enabling the extraction of tritium from the salt for storage in solid-state media.

Tritium Recovery & Permeation

Tritium recovery and permeation from molten fluoride salts in the molten salt breeder reactor (MSBR) has been previously studied, and is believed to be analogous to the application of such salts in the LIFE engine [Foster and Wright 1973; Moir et al 1984]. In the case of the MSBR, it was assumed that tritium was dissolved in the molten salt, with the formation of tritium fluoride (TF) suppressed. It was believed that tritium in the MSBR would permeate through the hot steel tubes of heat exchangers and steam generators at a rate of approximately 1 gram per day in the absence of permeation barriers, assuming that 1% of the helium coolant flow rate was processed for tritium recovery at 90% efficiency per pass (90% separation efficiency). Tritiated water in the steam system was considered to be a personnel hazard at concentration levels well below one part per million (< 1 ppm), a level that would be reached quickly without costly isotopic processing. The use of permeation barriers (coatings) on MSBR and steam generator tubes was

said to be able to reduce the leak rate of tritium into the steam system by two orders-of-magnitude (×100). For the option wi\th the lowest estimated leak rate of 55 curies per day, it may be possible to purge the steam system continuously to prevent the accumulation of tritiated water.

Structural Materials

The LIFE engine's structural challenges include: need for high-temperature strength; resistance to high-temperature creep; immunity to radiation damage, including swelling and helium embrittlement; resistance to corrosion and environmental cracking in high-temperature molten fluoride salts; and the ability to be fabricated into necessary shapes and configurations with practical welding processes. It is hoped that these challenges can be met with oxide dispersion strengthened (ODS) ferritic steels, sheets and coatings of refractory metals such as tungsten, advanced solid or liquid fuels, and other advanced reactor materials.

Specific requirements for the tungsten first wall are summarized in Table 5. This material will have to withstand temperatures of 650 to 750°C, and bombardment with 8.00×10^{22} neutrons per square centimeter per year (total of 3.20×10^{24} n/cm²), with 2.89×10^{22} neutrons per square centimeter per year (total of 1.16×10^{24} n/cm²) being fast neutrons. The corresponding damage will be approximately 36 displacements per atom per year (total of 1,440 dpa), with helium and hydrogen at 312 and 1,176 atomic parts per million per year (total of 12,480 and 47,040 appm), respectively. The ODS structural steel will have to withstand static and cyclic stresses, and corrosion by molten fluoride salts for 5 to 40 years, and may require protection with corrosion-resistant refractory-metal coatings.

The development of a LIFE engine for burning WG Pu and HEU requires materials of construction that can withstand harsh conditions of radiation, high temperature, active chemical agents, and pulsed (20/s) thermal and mechanical stress for periods of time as long as 10 years. Based upon preliminary investigations, it appears that tungsten armor could be used to protect the first wall, oxide dispersion-strengthened (ODS) ferritic steel could be used for fabrication of the structure; clad beryllium could be used as the neutron multiplier blanket; TRISO fuel could be used for the fission blanket; and graphite could be used as the reflector.

While these materials and materials systems are believed to be the best possible choices from the list of known materials, practical experience at the severe operating conditions expected in the LIFE engine does not exist. This system will require the use of molten fluoride salts at temperatures of 600 to 700° C, neutron doses of 3.5×10^{21} n/cm², with a resultant damage level of 150-200 dpa in the structural material, and 250-300 dpa in the solid fuel.

The service life of any construction material of will be limited by exposure to extreme temperature and radiation. Predictions of service life will require: radiation damage models, to enable accurate estimates of displacement damage (dpa), helium embrittlement (He/appm), hydrogen infusion (H/appm), and associated changes in physical and mechanical properties such

as changes in density (swelling); high-temperature phase stability, quantified through phase and time-temperature-transformation diagrams; high-temperature strength and creep resistance; corrosion resistance in high-temperature molten salts; static and cyclic stress analysis; and fracture mechanics, accounting for both the initiation and propagation of stress corrosion cracks, hydrogen induced cracks; and corrosion fatigue.

The corrosion and environmental fracture of the ODS steel engine chamber and fuel cladding in contact with the molten salt coolants and fuels must be fully understood. For example, it is believed that a chromium depletion layer will form in the 9-12Cr ODS ferritic steel, with a slowing of corrosion as the chromium concentration in the molten salt (Li₂BeF₄/PuF₃/UF₄) approaches saturation. Environmental cracking, including stress corrosion cracking, hydrogen (tritium) induced cracking, and corrosion fatigue, driven by the thermal pulsing at 10-20 Hz must also be understood. Ultimately, it will be necessary to predict the number of fatigue cycles possible, as a function of salt composition, temperature, stress amplitude, temperature, and neutron fluence.

The upper operating temperature limit of structural materials for reactors is determined by one of several factors, all of which become more pronounced with increasing exposure time, and which include: (1) thermal creep (grain boundary sliding or matrix-diffusion creep); (2) high temperature He embrittlement of grain boundaries; (3) cavity swelling (particularly important for SiC and Cu alloys); and (4) coolant compatibility/corrosion issues. In many cases, the upper temperature limit will be determined by coolant corrosion/compatibility rather than by thermal creep or radiation effects.

Broad ranges of high-melting radiation-tolerant structural material have been proposed for fusion-fission reactor structures. The properties of these materials are generally compromised by radiation damage. The upper operating temperature for each candidate structural material is summarized as follows:

•	Refractory Metals:	T < 4000°C
•	Carbides:	T < 4000°C
•	Nitrides:	T < 3500°C
•	Borides:	T < 3500°C
•	Oxides:	T < 3000°C
•	Intermetallics:	T < 3000°C
•	Silicides:	T < 2500°C
•	Sulfides:	T < 2500°C

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Nickel-Based Alloys (Hastelloys A, N, X; Inconels): T < 650-800°C

• Iron-Based Austenitic Steels (9-12% Ni): T < 600-650°C

• Other Steels (5-15% Cr): T < 500-600°C

• Ferritic Steels (3-3.5% Cr): T < 400-550°C

Metallic alloys that have been considered for structural components in variouss nuclear reactor applications include austenitic stainless steels, ferritic steels, and refractory metal alloys such as V-Cr-Ti, which are summarized in Tables 9 and 10. The austenitic steels are prone to extreme swelling and high temperature thermal creep, as will be discussed subsequently. Ferritic steels show relatively little swelling during neutron irradiation, and can be enhanced by inclusion of a nano-dispersion of oxide particles. The refractory may be useful at higher temperatures than the levels that can be attained with austenitic and ferritic steels. Refractory metals are found in Groups IVB and VB of the periodic table, and include Nb, Ta, Cr, Mo, and W, as serve as the basis of refractory alloys. Tungsten-based alloys and materials are frequently preferred for first-wall applications in fusion reactor designs.

Task 2.2 - Fuel Development

The primary objective of this portion of the proposed work is to develop a detailed understanding of the materials challenges for LIFE, the research and development plans for future work that enables the development of fuel forms. Two alternative fuel forms are being considered and evaluated for the denaturing and complete burning of WG-Pu and HEU: (1) TRISO fuels and (2) molten salt fuels

The relatively well-developed TRISO process is being favored at the present time for the production of fuels capable of surviving high burn-up and high neutron dose (fluence). Based upon a thorough review of the technical and scientific literature, it is believed that INL-based TRISO may already be a solid option for a 5-year weapons-grade plutonium (WG Pu) pebble capable of more than 99% burn-up in a LIFE reactor.

TRISO fuel fabricated directly from WG Pu and HEU would leverage the substantial advancements made by ORNL, INL and BWXT. Not that 79% FIMA (fraction of initial metal atoms) has been demonstrated with HEU TRISO fuel, with life limiting mode failures identified. At 79% FIMA, Kr-85 and other fission gas were released through cracks that formed in the SiC capsule, driven by fission gas pressure, as well as possible anisotropic swelling of the inner pyrolytic carbon layer. These failures have been analyzed, with steps formulated for further developing HEU and WG-Pu to even higher burn-up, with the ultimate goal being 95-99.9% FIMA. It is important to note that TRISO fuels operating in LIFE engines would be kept at a maximum temperature of 760°C, well below the 1100-1300°C levels proposed for applications in high-temperature gas-cooled reactors. Fortunately, this will (i) avoid the non-saturable void swelling regime for the silicon carbide capsule, and (ii) maintain the fuel operating temperature below the melting point of fission product silicides that could form, such as the well-known PdSi₅ compound. Other low-melting eutectics and solid solutions might also form, and must be investigated. All such interactions are being explored with thermochemical modeling using THERMOCALC, and database expanded with recent actinide data.

It is well known that pyrolytic carbons in TRISO can undergo anisotropic swelling along the aand c-axes. While continuous processing as done by the Germans substantially mitigates this problem, and is now being used in the United States, alternatives to graphitic layers and binder phases will also be explored.

Detailed finite element models of TRISO fuels have been developed with LLNL's ALE3D code, and have been successfully benchmarked with published IAEA standards for TRISO simulation. Thus far, sufficient detail has been incorporated into these models to account for SiC capsule wall stress due to fission gas accumulation inside the capsule, thermal pulsing due to laser firing, and de-bonding between layers. Work has also been done on thermochemical modeling of fission product interaction with the SiC capsule.

It would be particularly attractive to introduce WG-Pu or HEU directly into the LIFE engine, with as little pre-processing as possible.

One approach would be the direct digestion of WG-Pu and HEU into a high-temperature Li₂BeF₄ solvent/coolant as PuF₃ or UF₄, respectively. This digestion could be done well below the saturation concentration of either actinide fluoride salt. Here, too, detailed thermochemical modeling is essential. Such modeling will lead to a comprehensive understanding of actinide and fission-product solubility and any possible precipitation in such systems.

WG-Pu and HEU can be digested directly into a molten salt, thereby forming a high burn-up liquid fuel for the direct incineration of old weapons materials. Natural and depleted uranium (Nat-U and DU) can be incorporated into a variety of solid fuels including: TRI-structural ISO-tropic fuel (TRISO); baseball or solid hollow core (SHC); inert matrix fuel (IMF); and granular and powder fuels encapsulated in protective shells.

Baseline TRISO Fuel for LIFE Engine

Figure 15 shows an enhanced TRISO fuel, which has a more robust SiC capsule than conventional configurations to enable fission-gas containment at very high burn-up. This TRISO configuration is being considered as one possible high burn-up fuel option for the LIFE engine. The enhanced TRISO will have to endure more radiation damage than conventional TRISO fuels, as shown in Figure 16. The experience base for TRISO fuels is limited, with 8 to 20% FIMA demonstrated with low enriched uranium (LEU), and 79% FIMA demonstrated with highly enriched uranium (HEU) [Petti et al. 2004]. These LEU and HEU fuels experienced only approximately 3.5×10^{21} n/cm², but at a relatively high temperature of 1100 to 1200°C. In contrast, LIFE fuels will experience a much higher neutron dose of approximately 1.2×10^{23} n/cm², but at a much lower temperature of about 700°C. The temperature spike in the fuel pellets that result from the pulse of neutrons entering the fission blanket every 1/20th of a second is approximately 20°C.

Commercial TRISO fuels use a mixed ceramic phase, uranium oxy-carbide (UOC), which appear to minimize thermal migration effects of the kernel. Ultimately, several different types of fertile and fissile materials could be used for fabrication of solid fuels for the LIFE engine. Ultimately, solid fission fuel for the engine could be fabricated from nitrides, oxides, carbides, oxycarbides, silicides, aluminides and metallic alloys. The LIFE engine enable the direct burning of such fuels with no isotopic enrichment or chemical separation, with the potential to attain very high burnup, which is quantified in terms of fission of initial metal atoms (FIMA).

While the limitations of TRISO fuel at ultra-high burn-up are recognized and acknowledged, this fuel form is being used for the baseline design since the highest level of burn-up has been achieved with it. Based upon a thorough review of the technical and scientific literature, it is believed that an enhanced TRISO particle, based upon fuels already developed by INL, may provide an acceptable option for the nearly complete burning (> 99% FIMA) of weapons-grade

plutonium [Petti et al. 2004]. Other potentially superior fuels are being investigated in parallel, and include a new solid hollow core (SHC) fuel, inert matrix fuel (IMF), nano-structural fuels, powdered and granular fuels, and others.

In contrast, serious issues must be dealt with to achieve high-burn-up (> 99% FIMA) during 50 years of operation in a LIFE engine that is optimized to burn depleted uranium (DU). Challenges facing such enhanced high burn-up TRISO fuels in the LIFE Engine include: irradiation-induced damage of fuel materials, including swelling of the SiC and graphite; fuel failure due to the accumulation of noble fission gases such as Kr-85 within the SiC containment capsule; the thermo-mechanical response of non-isotropic PyC layers; which can be mitigated in part with uninterrupted TRISO processing; fatigue driven by thermal pulsing within the engine; thermal migration of kernel inside the SiC capsule, which can be mitigated in part by replacing UO₂ with UCO; attack the containment capsule by fission products such as Pd; and abrasion and corrosive attack of the fuel pebble. Table 7 summarizes these challenges, as well as possible mitigation strategies. The detailed research and development plan that will evolve in the future will lead to improved understanding through modeling and experimentation, and will provide the team solid guidance on exactly how to modify the well-established TRISO structure to withstand higher levels of radiation and fission gas buildup expected at such high burn-up.

LIFE TRISO fuel will operate below 1000°C, avoiding the non-saturable void-swelling regime. The fission blanket is expected to operate in the saturable point-defect swelling regime of silicon carbide (SiC), which is shown in Figure 17. Based upon published data for damage levels up to 8.5 displacements per atom (dpa), the swelling of SiC capsule expected to be 0.4 to 0.9 % [Snead et al. 2007]. Data at much higher irradiation levels is desperately needed, and will be generated during future irradiations with three-beam accelerators, high-flux hard-spectrum test reactors, and ultimately, LIFE pilot units. The three-beam accelerators will be used to simultaneously irradiate materials samples helium, hydrogen, and heavy ions to introduce damage.

Based upon a detailed kinetic model that accounts for the growth of dislocation loops, as well as ion-beam irradiation studies at Kyoto University's DuET facility, irradiation-induced swelling of SiC is expected to reach an asymptotic value of less than approximately one percent (~1%) as the dose increases [Ryazanov et al. 2002]. Model predictions and ion-beam irradiation of SiC show no significant swelling of above approximately 1 displacement per atom (1 dpa), as shown in Figure 18. However, the need to obtain data at higher neutron fluence cannot be overstated.

The anisotropic swelling of the inner and outer pyrolytic graphite (IPyC and OPyC) layers in TRISO particles, as well as the graphite binder in the pebbles, is considered to be a significant barrier to the development of ultra-high burn-up TRISO fuels, as shown in Figure 19. The swelling of the lattice normal to the hexagonal planes (along c-axis) is substantially greater than that parallel to the planes (along a-axis). A material that swells isotropically, and to a lesser extent, is preferable As shown in Figure 20, the fracture strength of irradiated nuclear graphite is expected to be 25 to 45 MPa at 20 to 25×10^{25} n/m², but may drop at higher doses [Zinkle 2008].

New Production Reactor (NPR) TRISO fuels were subjected to a neutron dose of 3.7×10^{25} n/m² (3.7×10^{21} n/cm²), and demonstrated the ability to achieve 64% when exposed neutrons from ATR, and the ability to achieve 79% FIMA prior to fission gas release when exposed to neutrons from HFIR. Differences observed between the experiments done at ATR and HFIR may be attributable in part to the softer neutron spectrum at HFIR. In regard to LIFE, additional testing and data are needed at doses as great as 10^{27} n/m² (10^{23} n/cm²) [Petti et al. 2004].

TRISO fabricated from HEU has achieved relatively high burn-up (79% FIMA) with our failure of the SiC capsule and leakage of fission gas such as Kr-85, as shown in Figure 21. The observed probability of failure appeared to be approximately 1 in 77,500 at this level of burn-up. Achieving such high burn-up with TRISO fabricated with natural or depleted uranium is a much greater challenge. Figure 22 shows fractures in the IPyC and SiC layers of a TRISO particle after reaching 79% FIMA. The observed leakage of fission gas from these particles appears to be due to the pressurization and fracture of the spherical silicon carbide pressure vessel.

One plausible explanation of the failure of the NPR TRISO at 64% FIMA is over pressurization of silicon carbide capsules with fission gas. The failure of these TRISO particles may coincide with the point where the SiC wall stress exceeds the ultimate tensile stress (UTS) of the material, as shown in Figure 23. Thus, one practical method of mitigating the failure may be to thicken the SiC wall.

Additional engineering of TRISO technology developed by DOE complex (INL, ORNL, BWXT and others) could lead to much higher burn-ups of natural and depleted uranium, with the mechanical ability to accommodate 95% to 99% FIMA. Enhanced TRISO particles have been designed with enough SiC wall thickness for fission-gas containment at 99.9% burn-up. Figure 24 shows a prediction of SiC wall stress verses fuel burn-up for this enhanced fuel. Crack initiation at defects in the SiC capsule must be accounted for in the development of enhanced TRISO fuels, and has now been modeled with finite element codes such as ALE3D and ABAQUS, as shown in Figure 25. The enhanced TRISO fuel will also experience dynamic stress due to thermal pulsing at 10 to 20 Hz in the LIFE Engine, which may cause thermal fatigue. Numerical analysis indicates fuel design will endure 10 to 20 Hz thermal pulse, as shown in Figure 26 [P. Damange, J. Marian, M. Serrano de Caro, LLNL, 2008]. Note that these models also incorporate the effects of radiation swelling on fuel materials; improvements in predictive ability over the coming years will increase confidence and lead to far better fuel designs.

Survival of the SiC capsule is complicated by reactions involving fission products such as Pd, which can be understood with the help of the phase diagram shown in Figure 27 [Okamoto 2000]. Deleterious low-melting eutectics form between fission products and encapsulation materials. A frequently cited example is the Pd-Si reaction. Pd₅Si phase formation threatens TRISO performance above the melting point of Pd₅Si, which is 836°C. Thermal migration of kernal in temperature gradient can bring such fission products in close proximity of the SiC. Mitigation includes the use of sacrificial SiC in (or around) the kernel and a ZrC diffusion

barrier. It is important to note that the TRISO operating temperature is expected to be below the melting point of deleterious phases such as Pd₅Si. Other possible phase transformations in the TRISO fuel are being predicted with THERMOCALC, and require knowledge of various binary pairs (C-Pu, O-Pu, O-U, Pu-U, Pd-Si, Pd-C, Si-C, etc.) as shown in Figure 28 [L. Kaufman, P. Turchi, LLNL, 2008].

The baseline TRISO fuel has several potential limitations. The mass fraction of fertile material in both conventional and enhanced TRISO fuel is limited by the packing efficiency of small TRISO particles (1-mm) in the larger TRISO pebble (2-cm), which is believed to be approximately 30%. The strength of the pressure boundary is limited by the properties of CVD silicon carbide. Furthermore, there is no means to control the buoyancy of the pebbles in the molten salt coolant.

Other Solid Fuel Forms

New fuel designs will enable us to overcome limitations of the current high burn-up solid fuels such as TRISO. The new solid hollow core (SHC) design being developed by LLNL may provide benefits unattainable with more conventional pebbles [Farmer, Latkowski and Abbott 2007-2008]. For example, higher mass fractions of fertile material can be incorporated into SHC than TRISO fuel. The predicted wall stresses in the SHC fuel at fission gas pressures corresponding to 20, 40, 60, 80 and 99.9% FIMA are shown in Figures 30 and 31, and are substantially less than those in comparable TRISO fuels [P. Damange, J. Marian, M. Serrano de Caro, LLNL, 2008]. The use of sacrificial SiC in the SHC should help mitigate attack of fuel materials by fission products. The specific use of sacrificial SiC in solid fuels is attributed to R. J. Lauf et al. [1984].

Fuel Fabrication Processes

The conceptual process for fabrication of enhanced TRISO fuel for LIFE from natural and depleted uranium is based upon conventional TRISO processing technology, and is shown in Figure 32. A process for the conversion of LWR spent nuclear fuel into IMF-type LIFE fuel is shown in Figure 33. Molten salt fuels do not experience radiation damage, and enable direct digestion of materials such as WG-Pu and HEU for incineration, and will be discussed in the following section entitled *Coolants and Liquid Fuels*.

Liquid Fuels & Continuous On-Line Reprocessing

If a homogenous liquid fuel is desired, fertile/fissile materials can be dissolved in these molten salt mixtures. Unlike solid fuels, molten salts with dissolved uranium, thorium and plutonium will not be structurally damaged by long-term exposure to neutron bombardment. From a purely technical point of view, such liquid fuels might therefore serve as ideal fuels for achieving nearly complete burn-up. Such molten salt fuels have been considered for graphite-moderated moltensalt breeder reactors that operate on the thorium-uranium fuel cycle, and enable continuous reprocessing of the fuel. Reprocessing is used to keep fission products at desirable levels. A

widely referenced fuel salt consists of Li₂BeF₄ (FLIBE) with dissolved fissile UF₄ and a blanket salt consisting of Li₂BeF₄ with dissolved fertile ThF₄. These conceptual designs assumed that NaBF₄ and NaF₂ would be used as secondary loop coolants. FLINABE and FLINAK based systems are also possible.

These liquid-phase solutions are very complex. In-depth understanding must be developed through application of internationally accepted predictive codes such as THERMOCALC to predict the formation of various phases, including precipitates such as plutonium trifluoride (PuF₃). The prediction of phase diagrams for complex liquid fuels, with large numbers of fission products, requires that the phase diagrams for several binary pairs of salts first be developed. For example, equilibrium phase diagrams have already been predicted for: BeF₂-LiF (FLIBE), BeF₂-ThF₄, BeF₂-UF₄, BeF₂-ZrF₄, LiF-PuF₃, LiF-ThF₄, LiF-ZrF₄, ThF₄-UF₄ and several other systems. The LIFE Project has recently predicted a phase diagram for the most basic FLIBE-based liquid fuel, assumed to consist primarily of lithium, beryllium, uranium and plutonium fluorides. This phase diagram is shown in Figure 35, and reveals regions of stable operation for such liquid fuels. Such systems would have to be operated at compositions and temperatures where no solid-phase precipitates form. Other continuous molten-salt processing schemes are reviewed elsewhere by Farmer, Shaw, Moir and Halsey [2008].

Most of our practical knowledge pertaining to liquid fuel comes from the development of the molten salt breeder reactor (MSBR) [Foster and Wright 1973]. The MSBR concept depends upon molten fluoride salts containing ^7LiF , BeF2, ThF4 and $^{233}\text{UF4}$ being pumped through a graphite moderated core where heat released by fission will raise the salt temperature to 700°C ($\sim 1300^{\circ}\text{F}$), close to the operating temperature of the LIFE engine. The heated salt then passes through an intermediate heat exchanger where its enthalpy is reduced by transfer of heat to a secondary salt, sodium fluoroborate, which in turn may transfer energy to steam at 3500 psia and 540°C ($\sim 1000^{\circ}\text{F}$).

Initially, a two-salt MSBR design was favored, with a fuel salt containing only fissile UF₄ and no fertile ThF₄, and a blanket salt containing only ThF₄ and no UF₄. The fuel and blanket salts were passed in close proximity to one another inside a graphite moderator block, where uranium-233 was bred from thorium. Unfortunately, problems due to dimensional instability of the graphite moderator block after long-term neutron irradiation were encountered.

In the case of the two-salt MSBR, a relatively simple on-line fuel reprocessing plant was enabled by the physical separation of fissile and bred uranium in the graphite moderator block [Figure 13.31, Foster and Wright 1973]. In the conceptual on-line salt reprocessing plant, fuel salt is fluorinated to convert dissolved UF₄ to gaseous UF₆, thereby allowing uranium separation from the liquid stream. The remaining salt is separated from rare earth and other fission products by vacuum distillation. Approximately 5% of the fuel salt is discarded in this conceptual process, with the need for LiF and BeF₂ makeup. In contrast, the blanket salt is fluorinated primarily for the removal of bred uranium from the fertile thorium. Approximately 5% of the blanket salt is

also discarded in this process, with the need for LiF, BeF₂ and ThF₄ makeup. The total salt leaving the loop is approximately 8%. A LIFE engine with liquid fuel would require a separation process for the continuous removal of rare earth fission products to prevent the precipitation of PuF₃. Similar salt removal would therefore be expected in a LIFE engine.

A reactor design with a single-salt two-region core and an easily replaceable graphite core assembly evolved, due to the problems anticipated with dimensional stability of the graphite in the two-salt design. This design eliminated the need for complicated flow paths inside the moderator block. The value of k_{∞} was found to peak at approximately 4% salt content and drop to below unity at approximately 18% salt content. It was found that a core with 13% salt content would be critical with k_{∞} of 1.034, while a reflector region having less than 37% salt content would be sub-critical with k_{∞} of 0.392. A breeding ratio of 1.07 to 1.08 was predicted for such a system with low fuel costs and small fuel inventories.

In the conceptual MSBR, the strong neutron capture of protactinium-233 ($\sigma_a = 43b$) in the core of the core leads to the production of protactinium-234, which in turn would decay to non-fissionable uranium-234. It is therefore be necessary to remove ²³³Pa (27.4-day half life) from the core so that it has time to decay to fissionable ²³³U. A liquid-to-liquid extraction process may be the best means for separating protactinium and uranium from the molten salt. An electrolysis process is then used to selectively recycle the uranium from salt in the decay tank to the MSBR fuel salt. Residual protactinium is then trapped in solidified salt for discarding.

This section of the report considers the destruction of WG-Pu by fissioning in a molten salt fuel. The solubility of plutonium and rare-earth fission products are both limited in these liquid fuels [L. Kaufman 2008]. However, with a feedstock of mostly ²³⁹Pu the reactivity of the fission blanket will be so high that a relatively small concentration of Pu will approach criticality, before its solubility limit will be reached. Much of the following discussion on molten salt fuels was written by a Livermore expert in molten salt breeder technology, Ralph Moir [2008].

Plutonium fuel has a high burn rate for a given fusion neutron flux of neutrons. Also based on prior studies, a high burn up (FIMA~70%) at relatively low neutron dose is expected, due to the high fission cross section, and the fission-to-total cross section of approximately 0.6. However, to burnout the higher actinides produced by the capture reactions, and to go to a burn up of over 99% FIMA, much higher neutron doses will be required.

The removal of noble gases such as xenon from the core of the MSBR is essential for efficient breeding. Specifically, helium purging is used to remove xenon-135 ($\sigma_a = 2.72 \times 10^6$ b), reducing the concentration of this gaseous fission poison by an order-of-magnitude (×10).

Plutonium oxide can be converted to plutonium tri-fluoride by hydro-fluorination at temperatures of 825 to 875K [Condit 2008].

$$PuO_2 + 3HF + \frac{1}{2}H_2 \rightarrow PuF_3 + 2H_2O$$

The PuF₃ is in granular form and can be added to the molten salt as discussed below. Uranium oxides can be similarly converted to fluorides. The digestion chemistry involves an extra step of the feedstock is in the form of metallic plutonium parts. Suitably small pieces would be feed into a furnace for hydride formation at 100 to 200°C.

$$Pu + H_2 + He \rightarrow PuH_2 + He$$

The PuH₂ is in a powdery form. Next the step is a conversion to fluoride at 100 °C or less in a nickel chamber. Helium aids removing the heat from the exothermic reaction:

$$PuH_3 + 3HF + He \rightarrow PuF_3 + \frac{3}{2}H_2 + He$$

The PuF₃ is in the form of fine grains. The forgoing is based on discussions with Condit [2008]. The above process steps would likely be a combination of batch and some continuous processes. Care would be needed to ensure criticality safety at all times under all conditions.

The granular PuF₃ can be augured slowly into a flowing FLIBE molten salt and allowed to dissolve or alternatively, crystalline BeF₂ and LiF can be added to the PuF₃ and then melt this mixture and add to the flowing FLIBE.

The practical operation of such liquid fuels depends upon the solubility of plutonium trifluoride (PuF3). The solubility of PuF3 is in turn strongly dependent on the lowest temperature of the molten salt in the circuit, and salt composition. As discussed elsewhere, the molten salt solvent is assumed to be FLIBE ($2\text{LiF} + \text{BeF}_2 \rightarrow \text{Li}_2\text{BeF}_4$). The solubility of PuF3 in this salt system with 33% BeF2 appears to be about 0.5 mol% at 600°C, as shown in Figure 36. The maximum temperature of materials containing the molten salt will ultimately be affected by the solubility limited lower temperature. The engine thermal-hydraulic design, mechanical-materials design and molten salt chemistry and solubility need to be further investigated.

Based on nuclear reaction and neutron transport calculations for a fission blanket populated with TRISO pebbles, engine designers believe that a mixture of FLIBE and 0.5 mole percent PuF₃ would behave similarly. The corresponding salt formulation is:

Formula: $0.663 \text{ LiF} + 0.332 \text{ BeF}_2 + 0.005 \text{ PuF}_3$

The effective mass density of plutonium at a plutonium concentration of 0.5 mole percent would be approximately 87 kg/m³. Neutronic calculations specifically tailored for a liquid fission blanket are in progress.

The solubility of plutonium and rare earth elements will be different in other molten salt mixtures. If engine designers find that they need higher plutonium concentrations than 0.5 mole percent, it may be possible to discover superior molten salt formulations by varying the BeF₂ content as suggested Figure 36. Additional data will be required to determine whether or not a solubility of approximately two mole percent plutonium tri-fluoride (~ 2 mol % PuF₃) can be achieved at 550°C. While extrapolations have been used to suggest that PuF₃ may have solubility >3.5% at 550°C, data at lower concentrations of BeF₂ are needed.

For the purposes of this discussion, it is assumed that the engine has a spherical blanket whose first wall is at 2.5 m radius, with a 0.15-meter beryllium pebble zone, extending outward to a radius of 2.65 meters, and a 0.5-meter molten-salt fission blanket, extending outward to a radius of 3.15 meters. This fission blanket may also contain graphite for moderation. In regard to the graphite content, two cases are assumed: 0 and 75 percent graphite. The corresponding volumes for the molten salt in these two cases are 53 and 13 cubic meters, respectively. Assuming a plutonium concentration of 0.5 mole percent, the plutonium concentration for these two cases would be 3750 and 940 kilograms, respectively. In the two cases it would take 3.75 and 0.94 years (always full power years) to burn-up one inventory of Pu and build up 0.18 mole percent of RE fission products.

As rare earth fission products accumulate in the molten salt fuel, the solubility of plutonium will diminish, and operation will become more difficult. To avoid the difficulties associated with growing concentrations of rare earth elements, and the precipitation of PuF₃ due to such growth, continuous removal of rare earths will probably be required. A conceptual process for removing RE fission products from LIFE liquid fuel is shown in Figure 37, and will also be discussed in context of removing noble metal fission products [Shaw 2008].

By fixing the inventory rare earth elements in the molten salt fuel, the required mass flow rate through the separation to achieve the assumed inventory can be calculated. Calculations indicate that a LIFE engine could have a continuous plutonium feed rate of 1000 kg/yr, with a flow rate through the on-line processing system of less than fifty cubic meters per year (50 m³/year) [Moir 2008]

Note that after 3.75 years, the initial inventory of plutonium will be burned and the concentration will grow to 0.18 mole percent. After burning 16 tons of plutonium in 60 years, the concentrations of Ba and Zr will be 2.2 and 2.9 mole percent, respectively, the concentration of RE fission products will be 2.9 mole percent, and the total concentration of all fission products will be nearly 9 mole percent. THERMOCALC predictions will need to be continued and include the following specific compositions:

Composition 1: $0.663 \text{ LiF} + 0.332 \text{ BeF}_2 + 0.005 \text{ PuF}_3$

Composition 2: $0.663 \text{ LiF} + 0.332 \text{ BeF}_2 + 0.005 \text{ PuF}_3 + 0.029 \text{ ZrF}_4 + 0.022 \text{ ZrF}_4$

It may be necessary to add uranium to the salt in order to minimize corrosion. In past work at ORNL keeping about 5% of the added uranium in the "+3" oxidation state was found desirable: UF₄/UF₃ ~ 20. The amount of uranium needed would likely be much less than the Pu and not affect the nuclear performance appreciably.

An estimate of the liquidus temperature from phase relationships in the LiF-BeF₂-PuF₃ system is approximately 600°C for a melt containing 0.5 mole percent PuF₃. The coldest part of the heat exchanger needs to be safely above the liquidus temperature. A 40°C margin would require 640°C. Assuming an additional 30°C drop across the boundary layer of the molten salt, the temperature at the inlet of the the molten salt blanket would be about 670°C. If the molten salt in the blanket is heated an additional 50°C due to fission, the blanket outlet temperature would be 720°C. If the hottest part of the blanket is cooled by the exiting molten salt with same boundary-layer drop as assumed before (30°C), then the structural hot spot temperature would be 750°C. If the hottest part of the structure is cooled with other than exiting FLIBE, this temperature would be lowered to 720°C. The engine wall temperatures are constrained by (Pu, REE)F₃ solubility considerations on the low end to structural materials temperatures limits on the high end.

It is believed that the LIFE engine will need considerable moderator in the molten salt region, probably graphite. This will greatly reduce the concentration of Pu at which fission will occur. His main trouble seems to be there are too many neutrons produced and no clear way to usefully use 'em. Options seem to be the addition of U, Th or Li. These will produce ²³⁹Pu, ²³³U or T, respectively. The production ²³⁹Pu in a ²³⁹Pu burner would be counter productive. Making ²³³U will have about 10% ²³²U resulting in gamma dose rates of many hundreds of Rem/h from 5 kg at 1 meter, and might therefore be a useful fuel feed for fission reactor that is proliferation resistant. Sales of both fuel and electricity and revenues from destruction of WG-Pu may enhance the economic viability of the system. The production of tritium could be used to fuel other LIFE engines whose purpose is to burn spent nuclear fuel.

Some fission products in a molten salt reactor are removed almost automatically and some can be removed by applying processes to side streams. Bubbles of inert gas, He, are injected into the molten salt flow. This serves to sparge gaseous fission products and including the volatile fluorides. In this way Kr and Xe are removed in an estimated time of 50 sⁱ. These gases are held up in pressurized vessels and allowed to decay.

The noble and semi-noble metal fission products do not form fluorides at the fluoride potential of the melt and are insoluble. They are: Zn, Ga, Ge, As, Nb, Mo, Tc, Ru, Rh, Pd, Ag, Cd, In, Sn, and Sb. They will plate out on the metal walls of blanket structures and piping and heat exchangers. If these elements circulate many times around the pumped circuit before plating out it may be possible to preferentially capture them on filters in a slipstream process loop in a time estimated at 2.4 h. One suggestion is to pass the molten salt through a nickel wire filter where plating occurs. Another suggestion is to use a centrifuge to enhance or concentrate these elements. This would be effective if precipitation is occurring homogeneously throughout the

molten salt. If, however, precipitation is occurring preferentially heterogeneously at metal surfaces then centrifugation would not be effective. These elements are then in metallic form ready to be converted to an appropriate waste form discussed in the next section.

The remaining fission product classes form stable fluorides and will circulate in the molten salt. They can be removed if we wish to add the necessary process equipment (reductive extraction). This process equipment allows removal of weapons useable materials and may not be desirable for LIFE from a proliferation point of view. In the case of burning weapon Pu this restriction probably would not apply.

As previously discussed, a side stream of molten salt can be contacted with a stream of liquid bismuth. A simplified flow sheet is shown in Figure 37. This process can remove noble and semi-noble metals, as well as Rb, Cs, Sr, Ba, yttrium, the rare earths, and zirconium. Estimates for removal times are 10 days for Zr, Rb and Cs and 25 days for the trivalent rare earths (Y, La, Ce, Pr, Nd, Pm, Gd, Tb, Dy, Ho, Er) and divalent rare earths and alkaline earths (Sm, Eu, Sr, Ba).

Task 2.3 - Structural Material Development

Iron-Based and Nickel-Based Steels

The performance of iron and nickel based alloys in nuclear applications is discussed in substantial detail by Ursu [1985, Section 12.5.2, p. 441-445]. Insights from Ursu are summarized in this section. The mechanical and corrosion performance, ease of fabrication, and relatively low cost of iron- and nickel-based steels make them logical choices for the fabrication of reactor structural components. The addition of refractory metals such chromium enhances high-temperature tensile strength, as well as corrosion resistance, though such performance is generally compromised by irradiation. The performance of iron and nickel based alloys in nuclear applications is discussed in substantial detail by Ursu [1985, Section 12.5.2, p. 441-445]. Insights from Ursu are summarized in several of the following paragraphs.

In the temperature range of 400-550°C, pearlitic-ferritic steels are frequently used. To extend the range of operating temperatures to 450-550°C, it is necessary to increase the chromium (Cr) content to 3-3.5 weight percent (3-3.5%). Carbide-forming elements such as molybdenum, tungsten, and vanadium (Mo, W, and V) are also beneficial for extending the range of operating temperature. These steels are said to behave acceptably with hydrogen within this range of temperatures.

Ferritic stainless steels are chromium-containing alloys with body-centered cubic (bcc) crystal structures [Davis 1995]. Chromium content is usually in the range of 10.5-30 weight percent (10.5-30%). Some grades contain molybdenum, silicon, aluminum, titanium, or niobium (Mo, Si, Al, Ti, or Nb) to confer particular characteristics. Ferritic alloys have good ductility and formability, but high-temperature strengths are relatively poor compared to those of austenitic grades. Toughness may be somewhat limited at low temperatures in heavy sections.

Annealed ferritic stainless steels usually have yield and tensile strengths of 240-380 MPa (35-55 ksi) and 415-585 MPa (60-85 ksi), respectively. Ductilities end to range from 20 to 35 percent. More highly alloyed ferritic stainless steels may have yield and tensile strengths as high as 515 MPa (75 ksi) and 655 MPa (95 ksi), respectively. These alloys are not strengthened by heat treatment or cold working. The strain-hardening rates of ferrite are relatively low and cold work lowers ductility.

Whereas martensitic stainless steels offer only moderate corrosion resistance, that of ferritic stainless steels can range from moderate for lower alloyed grades, to outstanding for more highly alloyed grades. Typical applications for ferritic stainless steels include equipment for handling oxidizing acids and organic acids, food processing equipment, and heat exchangers and piping used in seawater.

To enable operation at higher temperatures, in the 500-600°C range, additional chromium is required. Chromium additions of 5-15 weight percent (5-15%) increase the re-crystallization temperature and prevent plastic deformation. For example, enhanced steels with 12 weight percent chromium (12% Cr) have been shown to behave well at an applied stress of 200 MPa and a temperature of 500°C for 10.000 hours At the upper limit of this temperature range, molybdenum, tungsten, vanadium and niobium (Mo, W, V or Nb) are needed.

Austenitic steel with 18% Cr, 9% Ni and 1% Ti, with carbides, have been shown to be capable of withstanding an applied stress of 200 MPa at 630°C for 10,000 hour. Nickel alloys can be used as structural materials at temperatures up to 700°C, nickel-cobalt alloys can be used up to 800°C, and chromium alloys can be used up to 1100°C.

Refractory steels are used in the temperature range 900-1200°C, but cannot serve reinforcement purposes. Their strength depends upon temperature and increases with the rise in Cr concentration.

Oxide Dispersion Strengthened Ferritic Steels

New oxide dispersion strengthened (ODS) ferretic-martensitic (FM) steels have been developed specifically for a variety of nuclear applications, including cladding for fuel in fast reactors. These alloys are therefore considered to be good candidates for cladding for fuel in fast-flux reactors, as well as for the construction of LIFE engine structural components. These alloys exploit the ability of the ferritic matrix to withstand extreme neutron irradiation with relatively little swelling. Oxide dispersion strengthening with Y₂O₃ or TiO₂ nanoparticles enhances tensile strength and creep resistance at high temperature [Allen et al. 2005]. These alloys are therefore considered to be good candidates for construction of structural components of the LIFE engine.

The ODS FM steels known as Kobe 12YWT and 9Cr-ODS have demonstrated exceptional tensile strength and creep resistance at relatively high temperature, as shown in Figures 38 and 39 [Ukai and Fujiwara 2002; Klueh et al. 2007]. To further emphasize this point, the yield strengths and elongations as functions of test temperature for several additional ODS steels, including as-received EUROFER 97, CRPP ODS EUROFER97 with 0.3 wt. % Y₂O₃ and CRPP ODS EUROFER97 with 0.3 wt. % Y₂O₃ and 0.3 wt. % TiO₂ are shown in Figure 40 [Schaublin et al. 2006].

Given the thermal pulsing of the LIFE Engine, the resistance of materials of construction to thermal fatigue is also of interest. ODS FM steels appear to have much longer fatigue life than conventional ferritic steels such as modified 9Cr-1Mo steels, as shown in Figure 41. Predictions based upon regression analysis of data for $N_{\rm f} < 100,\!000$ cycles. Strain-controlled low-cycle fatigue tests were conducted for 9Cr-ODS and 12Cr-ODS at 873-1023K. The total strain ranges were controlled from 0.5 to 1.5% with a strain rate of 0.1% per second. Corresponding plastic strain ranged from 0.01 to 1%. The ODS has longer fatigue life than modified 9Cr-1Mo ferritic

steel. The oxide dispersion strengthening prevented softening or hardening during cycling [Ukai and Ohtsuka 2007].

Alternatives to Steel

Zinkle et al. have performed and outstanding critical analysis of the operating temperature windows for nine candidate structural materials for fusion reactors [Zinkle et al. 2004]. These include: four reduced-activation structural materials (oxide-dispersion-strengthened and ferritic/martensitic steels containing 8-12%Cr, V-4Cr-4Ti, and SiC/SiC composites), copper-base alloys (CuNiBe), tantalum-base alloys (e.g. Ta-8W-2Hf), niobium alloys (99Nb-1Zr), and molybdenum and tungsten alloys. The results are compared with the operating temperature limits for Type 316 austenitic stainless steel. Several factors define the allowable operating temperature window for structural alloys in a fusion reactor. The lower operating temperature limit in all body-centered cubic (bcc) and most face-centered cubic (fcc) alloys is determined by radiation embrittlement (decrease in fracture toughness), which is generally most pronounced for irradiation temperatures below similar to 0.3 T_{MP} where T_{MP} is the melting temperature. The lower operating temperature limit for SiC/SiC composites will likely be determined by radiation-induced thermal conductivity degradation, which becomes more pronounced in ceramics with decreasing temperature.

Vanadium Alloys

Vanadium alloys have been investigated for structural and first-wall applications in fusion reactors [Smith et al. 2000]. Insights from Smith et al. are summarized in this section. These refractory alloys provide favorable safety and environmental features such as low long-term activation, low decay heat and contact dose, and the potential for recycle. Vanadium alloys discussed by these investigators cover a broad range of compositions, V-(0-15)%Cr-(1-20)%Ti-(0-1)%Si, with particular emphasis given to a reference alloy, V-4Cr-4Ti. The three key alloying elements, vanadium, chromium and titanium, are mutually soluble in each other at elevated temperatures, and they all exhibit favorable low-activation characteristics.

Vanadium alloys with a few percent titanium are resistant to irradiation-induced swelling and embrittlement at projected operating temperatures, and are believed to be compatible with liquid-metal blanket/coolants. Helium and hydrogen transmutation rates in vanadium alloys are substantially less than those for other candidate materials exposed to high-energy neutrons characteristic of deuterium-tritium plasmas. Irradiation does harden the materials. For example, the yield stress of V-4Cr-Ti increases from the pre-irradiation level of 200-300 MPa to 600-700 MPa at a damage level of approximately 30 dpa, and temperatures of 430 to 600°C.

The primary thermo-mechanical treatment of these alloys is a solution anneal at approximately 1000° C for one hour. Properties are relatively insensitive to annealing conditions for 950-1100°C for 0.5-2 hours. Thermal aging at temperatures ranging from 500 to 1000C, for exposure times up to 5000 hours, show no significant grain growth (grain size $20 \pm 2 \mu m$). In summary,

candidate vanadium alloys exhibit good resistance to irradiation damage at temperatures of interest and offer a potential for long operating lifetime.

Other important physical properties of these candidate materials are summarized in Table 9 [Smith et al. 2000]. In particular, the coefficient of thermal expansion of V-Cr-Ti is lower than that of austenitic and ferritic steels, which would help alleviate thermal stress and associated environmental cracking, and the thermal conductivity is significantly higher, which would help alleviate concerns regarding heat transfer. This could make these materials good choices in applications where thermal pulsing is determined to be problematic.

Molybdenum Alloys

Molybdenum alloy with small additions of titanium, zirconium and carbon, such as 99.4%Mo-0.5%Ti-0.08%Zr-0.01%C, have good strain endurance and fracture toughness at high temperature. However, their susceptibility to oxidation requires that it be coated with thermal-spray coatings of Si-Al-Co. Other cladding is also possible. Additions of rhenium have been used to improve the ductility and fabricability of molybdenum alloys. Specific Re-containing molybdenum alloys that have been developed include: Mo-7Re (13 wt. % Re); Mo-27Re (42 wt. % Re); and Mo-30Re (45 wt. % Re). After neutron irradiation of Mo-27Re and Mo-30Re at 750 to 1075°C, MoRe₃ (Chi phase) precipitates formed at grain boundaries and on the surface at a damage level of 1.5 dpa; these precipitates were not observed in Mo-7Re [Ursu 1985]. Neutron activation may also pose challenges.

Niobium Alloys

These materials also have good strain endurance and fracture toughness. Some of the best tensile properties obtained with niobium-based alloys have been reported with: (1) Nb-15%W-3%Ti-1%Zr and (2) Nb-10%-3%Ti-1%Zr. Titanium additions have enhanced the hardness of these alloys, and have improved their refractory properties, allowing them to operate at temperatures as high as 1300°C. These materials tend to have better ductility than Mo alloys. Furthmore, these materials can be prepared using cold and hot pressing. Furthermore, argon arc welding is possible. Coatings of silicon have been used to prevent high temperature oxidation [Ursu 1985].

Other Advanced Materials of Interest

The LIFE engine will push crystalline engineering materials to their limits of radiation tolerance. The intense neutron bombardment will displace atoms from lattice sites numerous times, and may eventually convert crystalline materials into an amorphous state, with substantial helium and hydrogen implantation. Since a completely amorphous material may evolve at the extremes of neutron bombardment, the search for any new solid-phase material with immunity to extreme neutron irradiation should include amorphous materials (expected end state). Neutron bombardment would not displace atoms from lattice sites since none exist. The challenge in the development of radiation tolerant amorphous metals will be the attainment of high fracture

toughness, and incorporation of features that enable either trapping of large quantities of helium and hydrogen, or that enable the rapid transport of these species out of the matrix.

In addition to the possible application as radiation-tolerant structures, amorphous materials may also have other important applications in the LIFE engine. For example, the high boron content of Fe_{49.7}Cr_{17.7}Mn_{1.9}Mo_{7.4}W_{1.6}B_{15.2}C_{3.8}Si_{2.4} (SAM2X5) makes it an extremely effective neutron absorber for thermal neutrons, and suitable for criticality control applications [Farmer et al. 2007-2008]. Other yttrium-containing amorphous metals such as SAM1651 could be optimized for the absorption of fast neutrons by substituting Gd for Y or other alloying constituents, and should be explored for LIFE shielding.

Average measured values of the neutron absorption cross section in transmission (Σ_t) for Type 316L stainless steel, Alloy C-22, borated stainless steel, a Ni-Cr-Mo-Gd alloy, and SAM2X5 have been determined to be approximately 1.1, 1.3, 2.3, 3.8 and 7.1, respectively. This material and its parent alloy have been shown to maintain corrosion resistance up to the glass transition temperature, and to remain in the amorphous state after receiving relatively high neutron dose.

Materials used in criticality control applications must be relatively stable during irradiation with thermal neutrons. Melt-spun ribbon samples of various iron-based amorphous metals were subjected to high neutron dose in the 1.5 MW TRIGA reactor at McClellan Nuclear Radiation Center (MNRC) [22]. The neutron flux was 1.6×10^{10} n cm⁻² sec⁻¹. Samples were irradiated for three different times: duration of 1st irradiation was 44 minutes; duration of 2nd irradiation was 132 minutes; and duration of 3rd irradiation was 263 minutes. The corresponding neutron doses were: 4.3×10^{13} n cm⁻², 1.3×10^{14} n cm⁻² sec⁻¹ and 2.6×10^{14} n cm⁻², respectively. These doses are equivalent to approximately 670, 2000 and 4000 years inside the waste packages designed for emplacement at Yucca Mountain. An exposure corresponding to a 4000-year service life does not cause any detectable, deleterious phase transformations.

Predicting Degradation & Service Life of Engine Materials

Radiation Damage of Engine Materials

Effect for Radiation on Structural Steels, Refractory Alloys and Ceramics

Austenitic stainless steel (316), high-strength ferritic steel (HT-9), a refractory V-Cr-Ti alloy, and silicon-carbide, all of potential interest for application in the LIFE engine, have been irradiated with values of the radiation damage rate, helium transmutation rate, hydrogen transmutation rate, and nuclear heating rate measured. Irradiations were done with 14-MeV neutrons at a superficial power density of approximately 1 MW/m² for 1 year. These data are tabulated in Table 10 [Smith et al. 2000]. In general, ferritric stainless steels exhibit much less irradiation-induced swelling than Type 316 austenitic stainless steels, as shown in Figure 42 [Gelles 1996; Toloczko 2000; Klueh & Harries 2001; Zinkle 2008].

Annealed and aged ferritic stainless steel with fine niobium carbide (NbC) precipitates and relatively coarse Laves-phase particles was bombarded with a 4-MeV beam of iron ions. The damage level was 100 displacements per atom (100 dpa) at 400-600°C. Much of the damaged structure consisted of dislocation tangles that coarsened with increasing temperature. Cavity formation began at 450°C, appeared to peak at approximately 535°C, and was absent at 600°C. The maximum swelling observed during these tests was only 0.2 percent (0.2 %). The Laves particles exhibited recoil dissolution and restructuring [Farrell and Lee 1986].

Bombardments with iron, helium and deuterium (Fe, He, and D) beams or iron and helium (Fe and He) beams, at ratios of 10 and 45 atomic parts per million (appm) He/dpa, introduced high concentrations of small bubbles and comparatively few large bias-driven cavities. The latter were initiated at bubble clusters on niobium carbide (NbC) particles. The maximum extent of swelling was 0.75% and occurred at 550°C. Recoil dissolution of the Laves phase was retarded by the gases. Large critical cavity sized and an unbalanced dislocation and cavity sink strengths contributed to low swelling of this alloy. Such behavior makes dispersion strengthened ferritic steels generally attractive for LIFE applications.

Effect for Radiation on ODS Ferritic Steels

ODS ferritic steels have even better resistance to irradiation-induced damage than conventional ferritic steels without oxide dispersion strengthening. The alloy known as 9Cr-ODS was irradiated with Ni ions at 500°C and showed no detectable void formation at: 0, 5, 50, and 150 dpa, as shown in Figure 43. The effect of ion-beam irradiation on the oxide particles was also studied. The average size of these particles decreased from 11.7 nm for the un-irradiated case to 9.3 nm for the irradiated case at 150 dpa, as shown in Figure 44 [T. R. Allen et al. 2005]. Dispersions of nano-precipitates have also been shown to mitigate irradiation-induced swelling

attributable to helium, as shown in Figure 45. Helium appears to be trapped at nano-precipitates in the matrix [Mansur and Lee 1991].

ODS ferritic steel has been used successfully to fabricate prototype fuel pins for testing in Russia's BOR-60 reactor, a sodium-cooled fast reactor [Ukai et al. 2005]. No failures were observed at 5% burn-up (5% FIMA) and 25 displacements per atom (25 dpa). Tests are underway at this reactor which will provide data at 15% FIMA and 75 dpa. ODS alloys appear very resistant to irradiation effects, including high He levels, with a generally good balance of properties, and thus appear ideally suited for Chamber wall applications to 150 displacements per atom (150 dpa) and 1500 atomic parts per million helium (1500 appm He).

Friction stir, inertial and pressurized resistance provide best results Y_2O_3 particles remain in weld. Figure 46 shows a longitudinal cross-section in the vicinity of welded 9Cr-ODS steel cladding and its end-plug joined with pressurized resistance welding. Figure 47 shows fuel pins with ODS steel cladding and welded end-caps. Fuel pins with ODS steel cladding and welded end-caps, joined with pressurized resistance welding are being tested in BOR-60 reactor: 5% FIMA and 25 dpa were achieved in 2004; 15% FIMA and 75 dpa were achieved in 2008. To the best of our knowledge, no failures of the base metal or welds were experienced at 75 dpa [Ukai et al. 2005]. Others have also successfully joined ODS FM steels using industrial welding processes [Wright and O'Donnel].

Other approaches to dispersion-strengthening also exist, such as thermomechanical treatment (TMT). The microstructure of 9-12% Cr steels has been improved dramatically with hot rolling, which has become known as thermomechanical treatment (TMT). TMT has been used to process 9Cr-1Mo and 12 Cr steels, thereby forming precipitates at dislocations, as shown in Figure 48. Dispersions of precipitates produced with TMT are much finer than those observed in conventionally processed 9-12 Cr steels, and mechanical properties are substantially improved. [Klueh et al. 2005 & 2007].

Effect of Radiation on Nickel-Based Alloys

The irradiation induced swelling of austenitic nickel-based Inconel 600 and Type 316 stainless steel reflector components was studied in detail more than 20 years ago [Makenas et al. 1986, Table 1, Figure 11]. Transmission electron microscopy samples of Inconel 600 were prepared from two of the swelling samples used to generate the data in Table 11. Limited void swelling was found in the low neutron fluence specimen, while extensive voidage was found in the higher fluence, middle-of-core specimen. Irradiation-induced swelling of nickel-based austenitic steels such as Inconel 600 exceeds 5% at 9×10^{22} n cm⁻² and E > 0.1 MeV. Makenas et al. also discussed the implications of swelling on the performance of the reflector assemblies. Such swelling will probably make such austenitic nickel-based alloys unacceptable for widespread application in the LIFE engine.

The voids exhibit the truncated octahedral shapes normally found in face-centered cubic alloys. Neither transmission electron microscopy (TEM) nor the subsequent topical metallographic examination for second phases were helpful in determining the irradiation temperature of these specimens since Inconel 600 is not prone to temperature-dependent phase changes, which might indicate a specific temperature.

Effect of Radiation on Beryllium Neutron Multiplier

The general experience with beryllium (Be) is that it will swell and crack after receiving relatively low radiation doses, and that these problems will preclude its use for any structural function at doses as high as 200 dpa [Moir et al. 2007]. Beryllium swells by T and He produced during transmutation. At approximately 600°C, beryllium begins to release tritium (T), and continues the release up to 900°C. If beryllium is used only for non-structural components, such as pebbles or high-porosity metallic foams, with no requirements for strength and stability, problems associated with radiation-induced swelling may be inconsequential [A. Caro et al. 2007]. In summary, the integrity of Be components in the neutron multiplication blanket remains a worrisome problem that requires substantial attention.

Corrosive Attack of Engine Materials

Corrosion Mechanisms

The steels used for construction of the LIFE engine must be able to withstand high temperatures without melting or excessive thermal creep. Chromium has been added to many of the structural steels of interest to enhance both high-temperature mechanical properties, as well as oxidation and corrosion resistance. Table 12 provides some general guidance on acceptable ranges of composition for refractory oxidation-resistant steels. An alloy 0.35 to 0.45% for carbon, 2 to 3% for silicon, and 8 to 10% for chromium is suggested for operation at 850°C, close to the current design temperature of the LIFE Engine. The concentration of chromium in the ODS FM steels proposed for construction of the LIFE Engine fall within the recommended range.

In addition to having good high-temperature mechanical properties and being able to withstand oxidation, the structural materials used for construction of the LIFE Engine will have to withstand corrosive attack by the FLIBE (Li₂BeF₄) coolant. The corrosion and chemical compatibility of candidate metals and alloys is summarized in Table 13, entitled Materials Compatibility with Li₂BeF₄ Coolant, and is consistent with assessments in the literature [Ursu 1985, Section 12.4.4, Table 12.17, p. 436].

The corrosion of structural materials are due to the oxidative dissolution of reduced alloy constituents (M) with the high-temperature molten fluoride salt (*XF*), as described by Cheng et al. [2003]:

$$M + XF \rightarrow MF + X$$

In this reaction, *X* represents lithium, sodium, beryllium, or transmutation products, such as tritium. The transmutation of lithium produces hydrogen isotopes which react with free fluoride to form acid (*HF*, for example), making the blanket and coolant much more corrosive than it would otherwise be.

The relative corrosion susceptibility of refractory metals and Fe-Cr steels can be understood by ranking the free energies of formation for their respective fluoride salts, as done in Table 14 [Cheng et al. 2003]. The transmutation of lithium in the Li₂BeF₄ forms corrosive tritium fluoride, which behaves chemically like hydrogen fluoride, or hydrofluoric acid. The ultimate selection of materials for long-term containment such corrosive transmutation products in fluid pumping loops requires the use of alloy systems with constituents that form fluoride salts less thermodynamically stable than hydrogen fluoride. It is therefore useful to rank candidate alloying constituents based upon the free energies of formation of their respective fluoride salts, each with the units of kilocalories per gram mole of fluoride at 1000 Kelvin. Such a ranking reveals that MoF₆, WF₆, NiF₂, VF₅, VF₄ have less negative free energies than that of HF, indicating acceptability of Mo, W, Ni, and V, while FeF₂ has a free energy comparable to HF, indicating marginal acceptability of Fe-based alloys. Chromium has more negative free energy, which explains the dissolution of chromium by such fluoride salts, and the formation of the wellknown chromium depletion layer. Some refractory materials such as Nb-based and Ta-based alloys would be expected to be unacceptable for long-term exposure in high-temperature fluoride salts. Note that metallic beryllium (the proposed neutron multiplier in LIFE engine) would be expected to undergo dissolution in Li₂BeF₄, especially with galvanic coupling to more noble materials such as Mo-, W-, Ni-, V- and Fe-based alloys.

As previously discussed, molten salts used as blankets/coolants in fusion-fission hybrid reactors become extremely corrosive through the transmutation of lithium in the blanket/coolant, which results in the formation of hydrofluoric acid species (HF and TF). The corrosion mechanisms of steels and refractory alloys in these electrolytes are not yet completely understood, however, reasonable assumptions can be made based upon published experimental data. Corrosion may be controlled by manipulation of the oxidation potential, but it is expected that vigilant control would be required.

The postulated corrosion mechanisms for steels in molten fluoride salts are very complicated and are illustrated in Figure 49. The anodic process will involve the oxidative dissolution of metallic elements in the structural material, thereby forming dissolved metal cations. In the absence of oxygen, electrons generated by the anodic oxidation of the structural material must be consumed by some other electrochemical reduction process. Based upon published information, it is assumed that this cathodic reaction will involve reduction of H⁺ and T⁺ cations and the formation of gaseous hydrogen (H₂), deuterium (D₂), tritium (T₂) and other dimmer species formed from the three hydrogen isotopes. This reduction will liberate fluoride anions (F⁻) which will then form ion pairs with dissolved metal cations produced by the anodic dissolution of metallic elements from the structural materials. Some of these metal fluoride salts will be in

thermochemical equilibrium with volatile metal fluoride species. The high ionic strength of the molten salt makes it a highly conductive supporting electrolyte for corrosion reactions.

The published results of thermochemical simulation with the MALT2 database and the GEM code have provided valuable insight into the corrosion ferritic stainless steel and vanadium-based alloys in FLIBE [Nishimura et al. 2001]. Chemical species predicted to form in the gas phase include: (H,T)₂, (H,T)F, ((H,T)F)₂, (H,T)₂O, Be(H,T)₂, BeF₂, LiBeF₃ and FeF₃. Chemical species predicted to form in the condensed liquid phase due to the corrosion of ferritic stainless steel include: Fe, Cr, Fe₃W₂, Cr₂O₃, CrF₂, CrF₃, BeO, Li₂BeF₄ and LiBeF₃.

Experimental observations of the corroded surface were made with X-ray diffraction (XRD), Rutherford backscattering (RBS) and Moessbauer Spectroscopy have provided additional insight. The existence of α -Fe, Fe₂O₃ and spinel phases on the surface were detected with XRD. The formation of the chromium depletion layer after a 3-day immersion in the molten salt was found with Moessbauer Spectroscopy. Trivalent iron was also detected, suggesting the formation of Fe₂O₃. Rutherford backscattering (RBS) was used to measure the thickness of the oxygen penetration layer, which was found to be 200 nanometers.

These data indicate that the Cr₂O₃ film provides only limited corrosion protection. The preferential dissolution of chromium into the Li₂Be₄ salt results in the formation of a chromium depletion layer, which lies below a mixed oxide film of Fe(Fe,Cr)₂O₄ and disperse deposit of Fe₂O₃ particles. Dissolution and fluorination of iron leads to the formation of the volatile FeF₃ species. The formation of this volatile iron species can be suppressed by increasing the partial pressure of hydrogen partial pressure in the system. While the presence of metallic beryllium has a positive effect on the formation of corrosive HF (TF), it promotes formation of problematic gaseous BeH₂.

The transmutation of lithium in the Li_2BeF_4 salt leads to the formation of acidic tritium fluoride, which has the chemical behavior of hydrofluoric acid, as previously discussed. The reduction of T^+ at the ODS-salt interface may lead to the formation of reduced T_2 , with the simultaneous oxidation of Fe and Cr in the alloy to form Cr^{+3} and Fe^{+2} . Atomic tritium adsorbed on the surface of the ODS steel could be absorbed by the steel, which could lead to tritium (hydrogen) induced cracking. The presence of metallic beryllium is predicted to suppress the formation of HF (TF).

Cathodic tritium charging forms adsorbed tritium adatoms on the surface of the ODS FM steel. These tritium adatoms can recombine at the surface and form gaseous diatomic tritium. Alternatively, these adatoms may be absorbed into the bulk material and cause embrittlement and cracking, a situation that can be worsened by any dissolved species that might serve as a recombination poison. The mechanism would be analogous to that of hydrogen induced cracking (HIC). In many metals, such as titanium, infused hydrogen diffuses up stress gradients to crack tips, forming brittle hydride phases. These brittle hydride phases periodically rupture under the

stress at the crack tip, and enable crack propagation through the classical film rupture mechanism.

Furthermore, hydrogen isotopes absorbed by the bulk material may reduce oxide dispersion strengthening particles, further compromising high-temperature and radiation-tolerant mechanical properties. The possible reduction of oxide dispersion strengthening agents by absorbed hydrogen isotopes deserves further attention.

The localized corrosion, specifically the pitting of steels and refractory alloys has been reported in FLIBE molten salts. Pitting of alloys is frequently associated with the localized destabilization of passive films with halide anions, such as fluoride, the reaction of such halide anions with relatively soluble inclusions in the alloy surface, or the localized reaction of such aggressive anions with specific active sites on the surface of an immune metal. In all cases, a threshold electrochemical potential exists, above which localized attack can occur.

In experimental studies with vanadium, pits with diameters of approximately 200 microns were observed. After 10-day immersions, the alloy became brittle. X-ray diffraction did not show the formation of VF₂, VOF, or VO, which could have formed protective scales, but were not observed to do so. FLIBE is a very stable molten salt, however, under irradiation it forms acidic HF and TF, and causes severe corrosion of structural materials, including ferritic steels (specifically, Fe-9Cr-2W) and vanadium alloys (specifically, V-4Cr-4Ti). This does not necessarily rule out the design and fabrication of other high-performance ferritic and refractory alloys that could perform well in such environments. Alloys with elements forming less volatile fluoride species would be particularly interesting. Other pertinent corrosion studies have also been performed [Indacochea et al. 1999].

Corrosion Models

A wide range of vapor and aqueous phase corrosion models have been reviewed and documented by Farmer et al. [1991]. For example, crevice corrosion can occur in conventional light-water reactors, which leads to dramatic lowering of pH inside the crevice due to the hydrolysis of dissolved metal cations. Halide ions are simultaneously driven into the crevice by the potential difference that exists between the cathode at the crevice mouth, and the anode deep within the crevice. Low pH and high chloride ion concentration inside the crevice lead to accelerated corrosive attack, including other modes of localized attack such as pitting. Localized corrosion models include several for pit initiation: the halide nuclei theory; the point-defect model; and the stochastic probability model. Transport-limited models for the propagation pits are also reviewed, including cases where the growth of a hemispherical pit is limited by the presence of a salt film. The possibility of initiating a corrosion fatigue crack at pits is discussed, where the pit is considered to be a surface flaw. Models for initiation and propagation stress corrosion cracks via brittle film fracture at the crack tip are reviewed, and include the film rupture and slip dissolution models. Cases where the propagation of cracks is controlled by anodic dissolution at the tip, and assumed to be mass-transport limited are also considered. While relatively few

fundamental corrosion models have been successfully applied to the corrosive molten salt environment inside nuclear reaction systems, the possibility exists to do so. The successful corrosion engineering of the LIFE engine will makes such extensions of corrosion science essential.

Application of Mixed Potential Theory to Engine Materials

When two metals are galvanically coupled, a intermediate "mixed potential" is established. The exact position of this mixed potential, which is located between the oxidation-reduction potentials of the coupled metals, is determined by their relative electrode kinetics. The anodic and cathodic reactions are usually assumed to obey Tafel kinetics, and electro-neutrality must be preserved in the reactions. The intersection of the anodic and cathodic Tafel lines identifies the formal mixed potential. This will be the rest potential of both galvanically coupled metals. At this potential, one metal will undergo anodic dissolution, while the other will be cathodically protected, with the possible recombination of hydrogen ions on the surface. As an example, consider the zincs on your boats brass propeller, the less noble metal (zinc) will undergo anodic dissolution to protect the more noble metal from corrosion (brass or steel).

By galvanically coupling either metallic lithium or beryllium to iron immersed in a blanket of FLIBE, a mixed potential will be established for the couple between the two reduction-oxidation (redox) potentials. This mixed potential will be more positive than the reversible potential for either the lithium and beryllium couple and more negative than the reversible potential for either the iron and hydrogen couple. The lithium or beryllium will therefore undergo preferential dissolution, while the solid iron will be cathodically protected, and the hydrogen ions (from hydrofluoric acid species) will be converted to free fluoride anion and hydrogen gas. This may be an over simplified explanation, and the published standard potentials (which are directly related to the Gibbs free energies for the reactions) are for aqueous solutions (solutions with hydrogen ions). However, they do provide sufficient insight for a generalized understanding.

The coupling of a sacrificial material, such a lithium or beryllium, to a more noble material, such a ODS ferritric streel, serves to protect the steel at the expense of the sacrificial material. The extent of the protection provided by such galvanic coupling can be predicted through application of the mixed potential theory.

While the actual reactions involved in the corrosion of structural and cladding materials in the LIFE engine by molten salt coolants and fuels are numerous and complex, the following representative reactions can be used to shed light on the possible benefits of galvanically coupling beryllium and ODS ferritic steel. The anodic oxidation reactions for iron and chromium free electrons at the interface, and occur at nearly the same potential:

$$Fe \leftrightarrow Fe^{+2} + 2e^{-} - 0.409 \text{ volts}$$

$$Cr \leftrightarrow Cr^{+3} + 3e^{-}$$
 -0.410 volts

The transmutation of lithium in the molten salt will continuously generate a supply of tritium cations for the depolarization of the anodic corrosion reactions. Tritium cations are expected to behave electrochemically like hydrogen ions. The cathodic reduction of these dissolved cations will consume electrons produced during the anodic dissolution of the ODS steel.

$$2H^+ + 2e^- \leftrightarrow H_2 \quad 0 \text{ volts}$$

In this case, the dissolution rate of the engine wall will be bound by the tritium breeding rate, minus the rate at which tritium is removed from the salt. This reaction will serve as an ideal depolarization reaction for the anodic dissolution of iron and chromium, and may also provide adsorbed hydrogen isotopes at the interface which can diffuse into the bulk, thereby causing hydrogen embrittlement and hydrogen-induced cracking. Such bulk hydrogen could also reduce Y_2O_3 and TiO_2 nanoparticles, thereby removing the nanostructural feature responsible for high-temperature strengthening, and immunity to excessive irradiation-induced swelling. If this is found to be problematic, TMT-type materials may provide an opportunity for superior performance.

Beryllium can be galvanically coupled to the ODS steel, thereby enabling it to serve as a protective sacrificial anode.

$$Be \leftrightarrow Be^{+2} + 2e^{-}$$
 -1.70 volts

At the mixed potential that results from the coupling of the reactions for the anodic dissolution of iron and chromium, the reaction for the cathodic reduction of hydrogen isotope ions, and the reaction for the sacrificial anodic dissolution of beryllium, the anodic current densities and corrosion rates for iron and chromium will be suppressed, since these reactions will occur at a more cathodic potential than they would otherwise occur. The mixed potential can be substituted into the expressions for anodic current density, and the corresponding corrosion rate.

As shown in Figure 50, mixed potential theory applied to ODS ferritic steel without Be in molten Li₂BeF₄. In this case, the anodic Tafel line is assumed to be due to the anodic dissolution of iron and chromium from the ODS, while the cathodic Tafel line is assumed to be due to the electrolytic reduction of tritium cations. The intersection of these two Tafel lines defines the open circuit corrosion current, and the dissolution current density (corrosion rate).

As shown in Figure 51, mixed potential theory applied to ODS ferritic steel with sacrificial Be in molten Li₂BeF₄. In this case, the anodic Tafel line is assumed to be due to the anodic dissolution of sacrificial beryllium, as well as iron and chromium from the ODS, while the cathodic Tafel line is assumed to be due to the electrolytic reduction of tritium fluoride. The intersection of these two Tafel lines defines the open circuit corrosion current, and the dissolution current density (corrosion rate). Most of the anodic current density is attributed to dissolution of the sacrificial beryllium, which shifts the open circuit corrosion potential to more negative levels, where the dissolution rates of Fe and Cr are less.

As shown in Figure 52, mixed potential theory applied to ODS steel with and without beryllium in molten Li₂BeF₄. A comparison of these two curves, taken from Figures 50 and 51 respectively, clearly show the cathodic shift of the open circuit corrosion potential due to galvanic coupling of beryllium and ODS steel.

Cathodic Protection and Sacrificial Anodes

The corrosive attack of ODS FM steels in FLIBE may be manageable, provided that steps are taken to control the dissolution of the chromium into the fluoride salt. Corrosion resistance can be enhanced through careful engineering design, avoiding galvanic coupling of dissimilar metals, unless sacrificial protection is sought.

The use of sacrificial anodes is a form of passive electrochemical protection. Based upon preliminary analysis, it appears that either lithium (Li) or beryllium (Be) can be used as sacrificial anodes for the protection of the more noble ODS FM steel components. In the case of pure FLIBE, the introduction of metallic beryllium (Be) into FLIBE is considered a viable means of mitigating the corrosion of steels, since it promotes the conversion of hydrofluoric acid (HF or TF) species to hydrogen (H₂ or HT). If beryllium is used, the formation of colloidal beryllium in the molten salt may be problematic, in that it will thicken the solution. The formation of these particles may cause the salt solutions to become viscous, gray in appearance, and may necessitate continuous on-line removal. At this point, it should be noted that the beryllium neutron multiplier should be protected with a cladding or encapsulation to prevent unacceptable rates of dissolution and loss of dimensional tolerance during the service of the LIFE engine. Dimensional change would result in a loss of Be particle percolation in the annular regions of the LIFE engine, and jamming.

In essence, the beneficial effects observed by placing metallic beryllium in contact with neutron-irradiated FLIBE, with lithium transmutation products such as HF and TF can be understood, at least conceptually, by considering the electromotive series. Any electrochemical potential more negative (cathodic) than the stated Nernst potential will push the reactions to the right (reduction, acceptance of electron). Any electrochemical potential more positive (anodic) than the stated Nernst potential will push the reactions to the left (oxidation, liberation of electron by reduced metal or gas).

It is believed that metallic beryllium and lithium reduce hydrofluoric acid (HF/TF) formed from lithium transmutation products, and therefore eliminates aggressive free fluoride anions from the melt, along with the corrosive attack of structural materials caused by these species. The reactions for beryllium and lithium are:

$$2TF + Be \rightarrow BeF_2 + T_2$$
$$2TF + 2Li \rightarrow 2LiF + T_2$$

Since more LiF than BeF₂ is consumed within the fusion reactor, the use of metallic lithium is the preferred method of reducing hydrofluoric acid and replenishing the lithium in the moltensalt blanket and coolant. The dissolution of either can be actively controlled with an adder potentiostat if desired.

Similar electrochemical protection may be accomplished with adder potentiostats (operational amplifiers with formal reference electrodes). Instead of relying on a sacrificial anode to maintain the surface to be protected at the appropriate potential, the potentiostat can be used to apply the potential through application of a carefully controlled current. Thus, the redox state of the ferritic steel and transmuted lithium (H⁺ and T⁺) can be intentionally converted to the gaseous forms in a well controlled manner.

The ODS FM steel structure may also undergo corrosive attack by liquid fuels, which would consist of Li₂BeF₄ and dissolved UF₄, as well as a broad range of fission products. The ability of the broad range of fission products available at high burn-up to depolarize the anodic dissolution of the engine wall must be addressed. The use of sacrificial anodes and cathodic protection systems to mitigate the corrosion of the LIFE Engine in this aggressive environment should also be investigated.

An electrochemical controller could be used to mitigate the corrosive effects of HF and TF, and the attack of structural materials, such electrochemical control can also be used to manipulate the oxidation state of soluble uranium species (UF₃, UF₄ and UF₆). The formation of the volatile hexavalent uranium species should be avoided if possible.

The use of high-temperature cathodic protection systems to control corrosion and actinide volatility in liquid fuels should be investigated. Imposed currents could suppress corrosion current and convert any potentially volatile hexavalent uranium species to the less volatile tetravalent form.

In summary, the free energy of formation of fluorides of the alloying elements found in steels and refractory alloys can be used to determine the relative stability of these materials in molten salts. The reduction of lithium transmutation products (H⁺ and T⁺) drives the electrochemical corrosion process, and liberates aggressive fluoride ions that pair with ions formed from dissolved structural materials. Corrosion can be suppressed through the use of metallic Be and Li, though the molten salt becomes laden with colloidal suspensions of Be and Li corrosion products in the process. Alternatively, imposed currents and other high-temperature cathodic protection systems are envisioned for protection of the structural materials. This general approach is highly successful in the protection of ship hulls. Electrochemical control could prove to be enabling technology for such high-temperature molten-salt reactors.

Corrosion Rate Limited by Corrosion Product Solubility

Corrosion rates of ODS ferritic steel by Li₂BeF₄ and Li-Na-K-F coolants are expected to be less than approximately 1 mil per year. Corrosive attack by these coolants will probably be limited by saturation of the electrolyte with dissolved iron and chromium. The beneficial effects of Be on the corrosion of ODS can be understood by application of mixed potential theory. Several refractory metals are also appropriate choices for corrosion protection in high-temperature fluoride salts. ODS has been successfully joined using several different industrial welding processes, and provides exceptional high-temperature properties LIFE engine design

The corrosion of austenitic iron and nickel based alloys such as Type 316 stainless steel and Inconel have been studies by Oak Ridge National Laboratory in a variety of high-temperature fluoride salts, with the results given in Table 15. Results for the alloy INOR-8 are also reported. Corrosion rates for the iron-based stainless steel appear to be bounded between approximately 5 and 150 mils per year [Williams et al. 2006].

As previously discussed, chromium from the ODS ferritic steel is expected to dissolve into both the FLIBE in the primary cooling loop, as well as into the FLINAK in the secondary cooling loop. The dissolution of alloy chromium will be limited to some extent by the saturation concentration of the chromium in the fluoride salt. These limiting solubility values are given in Table 16, and have been used to estimate the maximum wall penetration expected in the steel LIFE engine. The tabulated solubility for beryllium in fluoride salts further illustrates the facile dissolution of the neutron multiplier in these salts. Uranium is of course also readily dissolved, which is beneficial in the formation of damage tolerant liquid fuels [Williams et al. 2006].

A simple model has been developed to account for the dissolution of LIFE Engine materials in the high temperature FLIBE and FLINAK. This model assumes a constant rate of corrosion until the solution becomes saturated with chromium. If the molten salt coolant or fuel saturates with iron and chromium corrosion products, the continued dissolution of the ODS ferritic steel is assumed to cease. The parameters used in this solubility-limited corrosion model for the LIFE engine are summarized as follows:

$$\rho_{ods} \approx 7.7808 \ g/cm^3$$

$$\rho_{salt} \approx 2.0000 \ g/cm^3$$

$$A_{ods} \approx 6,839,408 \ cm^2$$

$$V_{salt} \approx 1,248,053 \ cm^3$$

$$C_{salt,i}^* \approx 3,000 \ ppm \times \rho_{salt}$$

$$C_{salt,j}^* \approx 0.006 \ g/cm^3$$

$$r_{corr,j}^0 \approx 5 \text{ to } 71.1 \text{ mil/yr}$$

The solubility of chromium in various molten fluoride salts is summarized in Table 16. Limited corrosive attack of the ODS FM steels by the high-temperature molten fluoride salts has been predicted for two cases. Case 1 assumes a constant rate of attack of 5.0 mils per year, and cessation of corrosion at the point when the salt becomes saturated with dissolved chromium at a concentration of 3000 ppm as shown in Figure 53. Case 2 assumes a constant rate of attack of 70.1 mils per year, and cessation of corrosion at the point where the salt becomes saturated with dissolved chromium at a concentration of 3,000 ppm as shown in Figure 54.

Mechanical Fracture and Fatigue

The Griffith criterion for crack propagation relates the energy release rate to stress, flaw size and elastic modulus, and provides the theoretical basis for predicting mechanical fracture. This criterion enables a threshold (critical) stress to be calculated for a given flaw size, or vice versa [Thornton & Colangelo 1985, Chapter 9].

The effects of stress concentration at a crack tip are accounted for in a stress intensity factor (K_I) , which is calculated from the applied stress, characteristic crack dimensions, and modulus. Similarly, the threshold (critical) stress intensity factor (K_{IC}) is calculated from the critical stress, critical crack dimensions, and modulus. The stress intensity factor (K_I) for the concentration of tensile stress at a surface imperfection can be found in any one of several references [Andresen & Ford 1994; Farmer et al. 1991; Thornton & Colangelo 1985]. When the stress intensity factor exceeds the threshold $(K_I > K_{IC})$, the conditions for crack initiation and propagation are met.

Fatigue cracks may initiate and propagate when periodic fluctuations in stress are applied to a material. The fatigue crack velocity is proportional to the amplitude of the fluctuating stress intensity factor, raised to some power.

Environmental Fracture

When corrosive environments and tensile stress coexist, stress corrosion cracking (SCC) may occur. When SCC of a material occurs, the critical stress for fracture is lower than that for purely mechanical fracture. The straining at the crack tip causes slip and fracture of a brittle oxide film assumed to exist at the root of the crack. Once the passive oxide film fractures and the crack length is increased by an amount equivalent to the thickness of the film, the bare metal exposed at the crack tip is repassivation, and the film fracture process is repeated in a step wise manner.

Nakayama and Takano [1986] applied a slip-dissolution-repassivation (SDR) model to the SCC of Type 304 stainless steel in boiling 42% MgCl₂ solutions. Similar models have been developed for stress corrosion cracking and also assume that propagation is due to the fracture of the

passive film at the crack tip [Ford 1983; Andresen 1987-88; Jones 1987; Farmer et al. 1991]. Stress corrosion cracking (SCC) is assumed to initiate and propagate when the stress intensity factor exceeds the threshold stress intensity factor for SCC ($K_I > K_{IC}$). Once initiated, stress corrosion cracks have been shown to propagate at a rate proportional to the crack tip strain rate raised to the n^{th} power. The proportionality factor is also expressed in terms of the adjustable parameter n, which depends upon the repassivation kinetics at the root of the crack. The slip-dissolution model for stress corrosion cracking has been used to predict crack propagation in boiling-water reactors (BWRs) [Andresen and Ford 1994].

Other more fundamental models have been under development for more than two decades, but are not yet sufficiently mature to enable quantitative prediction of engine components. For example, a molecular dynamics (MD) model for the propagation of cracks in a surface covered with a thin brittle film was developed several years ago by Paskin et al. [1980-83]. Atoms were assumed to be arranged in a triangular lattice and to interact with a 6-12 Leonard Jones potential: Newton's laws of motion were solved, thereby enabling prediction molecular motion in the material near the incipient crack. An external tensile load was applied to each atom of the uppermost row, and each atom of the lowermost row of the sample. While such simulations provide invaluable insight into the mechanisms of SCC and other environmental cracking, quantitative predictions in realistic environments may not yet be possible for the LIFE engine, and more empirical models may still be required. More recently, others have done additional work on simulating environmental cracking of alloys, including SCC, with MD techniques.

End of Fuel Cycle Implications

The end of the fuel cycle for DU and WG-Pu burning LIFE engines are similar, since the DU burning machine actually burns plutonium during most of its period of operation. As both machines approach 99% burn-up, the high-level radioactive wastes for the two systems become similar in composition and other characteristics, and will require similar geological disposal. Both machines generate far more electricity per mass of nuclear waste generated that light water reactors. This section discusses issues pertaining with the disposition of a spent LIFE fuel after nearly all of the plutonium is converted to fission products.

The actual worldwide SNF inventories from the existing fleet of light-water reactors, and the corresponding number of Yucca Mountain Equivalent (YME) repositories is summarized in Table 17. The equivalent worldwide SNF inventories from a hypothetical fleet of LIFE engines with generating capacity comparable to the existing fleet of light-water reactors, and the corresponding number of YME repositories is summarized in Table 18. The SNF generation and required YME repositories for the actual LWR fleet, and the hypothetical fleet of LIFE engines is shown in Table 19. The YME repository is assumed to have a disposal capacity of approximately 70,000 MTHM (capacity of 63,000 MTHM reserved for commercial fuel).

Estimates in Tables 17 through 19 are based upon data for the worldwide fleet of light water reactors, and the spent nuclear fuel generated by those reactors, and was taken from a recent publication by the International Atomic Energy Agency [Fukuda et al. 2007]. The assumed cost for a Yucca Mountain Equivalent (YME) repository was based upon recent statements by the United States Department of Energy [Nuclear Waste News 2008]. The Cost for the Yucca Mountain Repository has increased from the original life-cycle cost of \$58 billion to more than \$90 billion. The increased cost is due to a variety of factors, including the cost of temporarily storing the current of spent nuclear fuel, as well as escalating costs for construction and procurement of materials for the engineered barrier system. Given the escalating cost for geological disposal and such strong public and governmental opposition, any nuclear power-generation that can increase the quantity of electrical energy serviced by facility such as Yucca Mountain, thereby decreasing the need for a larger number of future repositories, should be welcome.

The mass flows and transuranic waste generation by various fuel cycles, including two based upon LIFE engines, are compared in Table 20. The data for the various non-LIFE fuel cycles was taken from publications by the NEA [Advanced Nuclear Fuel Cycles and Radioactive Waste Management, NEA No. 5990, Nuclear Energy Agency, Organization for Economic Co-Operation and Development, 2006, p. 30, Table 2.2].

The LIFE engine's once-through fuel-cycle story is summarized in Figure 55, and includes breeding and incineration in the LIFE engine, interim storage of the spent fuel removed from the engine, and the ultimate disposal of that waste in a geological repository [Farmer et al. 2008].

First, natural uranium, depleted uranium (DU), light water reactor spent nuclear fuel (LWR SNF), highly enriched uranium (HEU), or weapons grade plutonium (WG-Pu) is burned in the LIFE engine to 80 to 99.9% FIMA. The inventory of transuranics (TRU) in the fission blanket of LIFE engine at various levels of burn-up is shown in Figures 56 and 57 [Shaw et al. 2008].

Assuming current statutory limits for the disposal of spent nuclear fuel, a single YME repository could service more LIFE engines that light-water reactors. LIFE could extend the useful service life of deep geologic repositories by factor of twenty (×20). LIFE would require less repository capacity than LWR. Twenty times more electricity could be produced with nuclear before filling up repository (Figure 58). A single YME repository could enable more electrical power generation with a fleet of LIFE engines than with a fleet of light-water reactors. LIFE could extend the useful service life of deep geologic repositories by factor of twenty (×20) (Figure 59). These calculations assume that each reactor will have a service life of approximately 50 years: LWR = 1.2 GWe per 900 metric tons of heavy metal waste; LIFE = 1.0 GWe per 40 metric tons of heavy metal (MTHM) waste [Farmer et al. 2008].

A more complex scenario for the transition from the current fleet of light water reactors to LIFE engines is shown in Figures 60 through 61. Electrical generating capacity for the first scenario is shown in Figure 60. In this case, it is assumed that the LWR fleet grows so that it can continue to provide 16% of the projected demand, and that no LIFE engines will be built. This scenario leads to a relatively large energy deficit, with the demand far exceeding the available electrical power. Electrical generating capacity for the second scenario is shown in Figure 61. In this case, it is assumed that LIFE engines are introduced in 2030 to fill the projected need for electrical power. The total SNF that would have to go to a geological repository for permanent disposal for the two scenarios is shown in Figure 62. In the second scenario, it is assumed that the LWR SNF that is generated would be burned in the emerging fleet of LIFE engines. The number of YME repositories for the two scenarios is shown in Figure 63 [Farmer et al. 2008].

Limited Production of Weapons-Attractive Materials

LIFE engines will produce less transuranic (TRU) and plutonium (Pu) for immobilization and disposal than comparable light water reactors. A LIFE engine's more complete burn promises to generate far less plutonium per reactor than typical LWR (Figure 64). A LIFE engine with an assumed electrical generating capacity of 1 GWe will discharge less than one kilogram of plutonium (1 kg Pu) after 99.5% FIMA. In contrast, an LWR discharges approximately 9,000 kilograms of plutonium (~ 9,000 kg Pu). The more complete burn possible with a LIFE engine will also generate far less TRU than a comparable LWR (Figure 65) [Farmer et al. 2008].

The smallest unit of fission fuel from the conceptual LIFE engine is a spherical pebble, and is believed to be far less attractive for possible theft than the smallest unit of fission fuel from a typical LWR, which is a single cylindrical rod. At the end-of-burn, a fuel pebble from a LIFE engine has a calculated specific activity of approximately 2.1×10^6 Ci/kg-TRU, compared to a

specific activity for fuel rod from a typical LWR of approximately 2.3×10^4 Ci/kg-TRU. Fuel pebbles removed from the LIFE engine will contain much less plutonium than fuel rods removed from a typical LWR, but will be much hotter, both thermally and radioactively. A single fuel pebble from a LIFE engine will therefore be a far less attractive target for theft than a single fuel rod from a typical LWR (Figures 66 and 67) [Farmer and Zhao 2008].

Interim Storage

After removal from the LIFE engine, the fuels will be thermally and radioactively hot, and will require cooling to survive in interim storage and the repository. Prior to packaging, LIFE pebbles could be initially cooled for several months to five years in a relatively high temperature cooling fluid such as FLINAK. Once the pebbles are packaged, the surface of the package must be kept cool during interim storage, transport and eventual emplacement underground.

The pseudo steady-state temperature profile inside a solid fuel inside a metal container can be described quantitatively by combining two well-known solutions to the transport equations for heat transfer found in Theodore's book on transport phenomena [Theodore 1971]. The result of combining equations from the aforementioned examples enables calculation of the maximum center-line temperature of the fuel mass inside the container and is summarized as follow:

$$T_{\text{max}} = T_{od} + \left\{ \frac{q_{r=a}}{k_{metal}} \right\} \times \left\{ a \ln \left(\frac{a}{b} \right) \right\} + \left\{ \frac{Ab^2}{4k_{fuel}} \right\}$$

The quantity $q_{r=a}$ is the heat flux on the outer surface of the container, which is calculated from the thermal output of the fuel at any point in time, k_{metal} is the effective thermal conductivity of the metal container, A is the power density in the volume occupied by the fuel, k_{fuel} is the effective thermal conductivity of the fuel mass, a is the outer radius of the container, and b is the outer radius of the fuel mass (inner radius of container). Since the fuel mass consists of TRISO pebbles, with conductive filler in interstitial volumes between pebbles, the fuel thermal conductivity is degraded by multiplying it a quantity defined as the *area fraction available for heat conduction* (f). The effective fuel thermal conductivity is then calculated as:

$$k_{fuel} = f \times k_{fuel}^{0}$$

The calculation accounts for an additional heat transfer resistance at the surface of the container. The flux of heat into the surrounding air is:

$$q_{od} = h_{air} (T_{od} - T_{air})$$

Reasonable ranges for the heat transfer coefficient range from 3-11 W/m²-K for natural convection in air, and 28-114 W/m²-K for forced convection in air [Bennett and Myers 1974]. Values of thermal conductivity are for silicon carbide (fuel) and nickel-based container material. Quantities are further defined in Table 21 [Farmer et al. 2008].

The characteristics of a hypothetical interim storage facility for LIFE spent nuclear fuel (SNF) are summarized in Table 22 and 23 [Farmer et al. 2008]. The spent TRISO fuel from a single LIFE engine, assuming that the machine was fueled with 40 metric tons of depleted uranium, is contained in 10 to 11 containers, each with a length of 5.95 meters. These containers are assumed to sit horizontally in a square flow channel (4.2 m × 4.2 m) with a closed-cooling air system. A cooling air velocity of 1 meter per second corresponds to a volumetric flow rate of 695 cubic meters per minute, with an assumed entrance temperature of 25°C, and an exit temperature of 60°C. The calculated maximum temperature inside the fuel mass is 261°C for an assumed heat transfer coefficient of 11 W/m²-K and an *f*-value of 0.10, and 359°C for an assumed heat transfer coefficient of 11 W/m²-K and an *f*-value of 0.05, both of which are well within comfortable operation for the TRISO fuel. In both cases, the surface temperature of the metal container is about 151°C, and the temperatures at the fuel-container interface is about 163°C, well below the temperature where intermetallic phases deleterious to long-term corrosion performance precipitate in Ni-Cr-Mo alloys such as Alloy C-22.

The temperature levels that will be experienced by the fuel materials during interim storage have been calculated, with the results also summarized for two different heat transfer scenarios in Tables 22 and 23 [Farmer et al. 2008]. With proper ventilation, the centerline of the stored spent fuel from the LIFE engine should not exceed 250 to 300°C. This temperature is well below the melting points of fuel materials, as well as the melting points of eutectics that are expected to form between silicon carbide encapsulation materials and fission products such as palladium. The Repository Design Team for LIFE continues to explore and optimize the interim storage and repository design for this hybrid fuel. The predicted power, cooling air flow, and temperature for a container filled with LIFE spent fuel during dry interim storage are shown in Figures 68 and 69, respectively [Farmer et al. 2008]. The close-packed TRISO pebbles inside the container are assumed to have an effective fractional area for thermal conduction of 0.10 (factor used to degrade thermal conductivity to account for interstitial spaces). After 100-years in such interim storage, the centerline temperature of the fuel mass should decay to a temperature corresponding roughly to the boiling point of water.

Geological Disposal of Spent LIFE Fuels

Following the initial cooling at the reactor site, a geological repository similar to Yucca Mountain (YM) may be used for long-term storage or disposal. The size of a geological repository to accommodate a entire fleet of LIFE engines (with the same generating capacity as our current light water reactor (LWR) fleet with a once-through fuel cycle) will be only 5 to 10 percent (approximately 7%) of that required for disposal of LWR spent nuclear fuel (SNF) in a geological repository similar to YM.

An Artist's rendering of a drift inside the YM geological repository with temperature limits is shown Figure 70. When completed, the repository will have 108 parallel drifts with a centerline spacing of approximately 81 meters. The drift wall temperature must be kept below 200°C to

minimize deleterious mineral transformations and swelling. The mid-pillar temperature, at a distance of 40.5 meters from the drift centerline, must be kept below 96°C to permit drainage of the infiltrating water. Next, the temperature of the drift walls must remain below 200° C to minimize swelling of the rock due to mineral phase transitions. Finally, the mid-pillar temperature must remain below 96° C (water boiling temperature) to permit drainage of infiltrating water between drifts.

Waste packages filled with spent fuel from LIFE satisfy repository temperature limits. The container, which will be fabricated from austenitic nickel-based Alloy C-22 must be maintained below 300°C to minimize formation of deleterious (P, σ and μ) intermetallic phases which deplete the matrix of those alloying constituents (Cr, Mo and W) which impart outstanding corrosion resistance. In the case of LWR SNF, the temperature limit for the zirconium alloy cladding is approximately 350°C. In the case of LIFE SNF, this would be replaced by the upper temperature limit for the TRISO fuel, which is assumed to lie between approximately 700 and 1400°C. Consistent with the current license application for YM the waste package outer shell temperature must remain below 300° C to minimize deleterious phase transformations in the Alloy-22 (precipitation of P, σ and μ phases), which deplete the matrix of those alloying elements responsible for the containers long-term corrosion performance.

Sufficient air flow through the interim storage facility and repository drifts will be required to ensure that the three temperature limits for the repository are not exceeded during the storage or disposal of LIFE SNF. The following assumptions were made to enable temperatures inside the repository to be predicted: (1) LIFE engines fueled with 40 metric tons depleted uranium; (2) incorporation of this uranium into the kernels of 1-millimeter diameter TRISO particles; (3) the incorporation of these small TRISO particles into approximately 13-14 million 2-centimeter TRISO pebbles; (4) operation at full electrical power of 779 megawatts for 40 years; (5) operation at a reduced power level for 10 years to incinerate the remaining actinides, with an ultimate burn-up of 99% fraction initial metal atoms (FIMA); (6) post-incineration aging of the TRISO pebbles for 5 years prior to emplacement in the underground interim storage facility; (7) post-aging specific thermal power of 548 kilowatts per 40 metric tons of burned depleted uranium, based upon ORIGIN2 simulation; (8) placement of the 13-14 million TRISO pebbles into 10-11 standard spent nuclear fuel containers, adapted to accommodate the spherical fuel pebbles at a sub-theoretical packing fraction of 60 percent; (9) each container will have an inner length of 4.87 meters, an outer length of 5.95 meters, an inner diameter of 1.64 meters, an outer diameter of 2.72 meters, and a volume of 10.26 cubic meters; (10) an end-to-end separation of 10 centimeters; (11) an initial linear power density of 8.8 kilowatts per meter after post-incineration aging of 5 years; (12) a linear power density of 1.45 kilowatts per meter inside the geological repository; (13) a thermal conductivity inside the waste package of 87 watts per meter per Kelvin, and (14) a pebble centerline temperature of less than 900°C.

A comparison of the heat generated by containers filled with LIFE and LWR spent nuclear fuels during the first million years is shown in Figure 71. Thermal power of LIFE SNF decreases with increasing burn-up (%FIMA). The difference in FIMA power is burnup of actinides; fission products are in quasi-equilibrium. The higher LIFE thermal power can be accommodated by repository design options. Linear power density in the repository (kW/m) decreases with increasing burn-up (%FIMA) [Blink et al. 2008].

Assuming the most conservative case, the maximum possible power density (closest packing) of the spent LIFE fuel and no cooling during interim storage at the LIFE engine site, the required cooling air velocities to meet these operational temperature limits during such long-term storage or disposal in an underground facility similar to YM are approximately 2.6 feet per second. This cooling air velocity, while well within engineering bounds, can be lowered by distributing the spent LIFE fuel over a larger area. If interim storage is used to cool the fuel, the power of each spent fuel container drops substantially. Consequently, the required ventilation air velocity in a geological repository would also drop, with an expected velocity of ventilation in the repository comparable to that expected for a repository filled with containers of LWR spent nuclear fuel.

Ventilation will be used to keep the temperature of LIFE SNF containers within limits during the pre-closure period of the repository, in a manner analogous to that for LWR SNF, as illustrated in Figure 72. This prediction of LIFE SNF container (waste package) temperature is for containers located near the drift's ventilation air exit. The ventilation air flow is assumed to be approximately 15 cubic meters per second (15 m³/s), with a velocity in the drifts of approximately 1.79 miles per hour (1.79 mph) or 2.6 feet per second. In addition to air flow, emplacement sequencing is also used as a means to control repository temperature within acceptable bounds. For example, in the case discussed here, it is assumed that all LIFE fuel is located in a centralized underground interim storage facility with the length of a single emplacement tunnel (drift) being 666 meters, which is long enough to accommodate 112 containers. Emplacement would be done sequentially, with seven 16-container groups moved into the drift at 10-year intervals. The saw-tooth pattern in the temperature-time profile is due to the phased, sequential periodic emplacement of the containers from the LIFE engine every 10 years. In this particular case, the burn-up is assumed to be approximately 99% FIMA [Blink et al. 2008].

The radioactivity of LIFE SNF has also been modeled and decreases as the burn-up (%FIMA) increases, as shown in Figure 73. The difference in FIMA power is burn-up of actinides; fission products are in quasi-equilibrium. Estimates for LIFE include some radionuclides (RNs) not included in the available estimates for LWR SNF, also called commercial spent nuclear fuel (CSNF). For example, Cm-246 is more important in LIFE SNF than in LWR SNF due to the high neutron fluence in LIFE (8 n captures). Other RNs such as Zr-93 and Nb-93m need to be closely scrutinized to determine why they are not included in available estimates for CSNF [Blink et al. 2008].

In one LIFE plant simulation, after approximately 40 years in the LIFE reactor, the activity of a single 2-centimeter diameter TRISO pebble is estimated to be significantly less than 400 curies, and comparable to the fuel rods removed from light-water reactors. After aging outside the LIFE reactor in interim storage, the activity decays relatively quickly. After aging for 10 years, the activity of a single pebble is predicted to drop by more than one order-of-magnitude to less than 15 curies. Then, after aging for an additional 10 years, the activity drops below well below 10 curies. After 50 years (comparable to period required retrieval of LWR fuel in YM), the activity drops to well below 5 curies, and after 100 years, the activity is well below 1 curie.

While the radioactivity and thermal output are comparable to current fuel forms, the criticality safety for any high burn-up TRISO fuel may be appreciably better than LWR fuel assemblies. This waste form may provide substantial benefits in during disruptive volcanic and seismic events. The fission products are contained in the pebbles and silicon carbide capsules, which will help isolate the waste from the surrounding environment. Current waste packages rely to some extent on borated stainless steel basket assemblies for criticality control within the containers. Such reliance would be much less important in a distributed fuel form such as TRISO or IMF.

Key radionuclides (RNs) identified as dominating the doses for different time periods in the YM repository's total system performance assessment for license application (TSPA-LA) are summarized in Table 24 [Shaw et al. 2008].

A comparison of energy-normalized inventories of RNs considered significant for the performance of the YM repository is shown in Figure 74. Plotted is the maximum ratio (at any time) of the activities per unit electrical energy generated by spent LIFE fuel (DU, 99% FIMA) and CSNF ($41.2 \text{ GWt-day/MT} \sim 4.3\% \text{ FIMA}$) [Shaw et al. 2008].

Similarly, a comparison of mass-normalized inventories of RNs considered significant for the performance of the YM repository is shown in Figure 75. Plotted is the maximum ratio (at any time) of the specific activities (Ci/MTIHM) of spent LIFE fuel (DU, 99% FIMA) and CSNF (41.2 GWt-day/MT ~ 4.3% FIMA). The most significant nuclides are less abundant in LIFE fuel, but several are higher. Some nuclides in LIFE fuel are not accounted for in the Yucca Mtn. TSPA (93Zr, 244Pu, 246Cm, 248Cm) [Shaw et al. 2008].

The risk-to-benefit ratio for a YME repository completely filled with spent nuclear fuel from a fleet of LIFE engines compared to that of a similar repository completely filled with spent fuel from a fleet of light water reactors, showing risk for the LIFE scenario, is shown in Figure 76. The dose predicted by performance assessment that is attributable to the generation of a given quantity of electricity is approximately ten times (~10×) less for LIFE waste than for LWR SNF [Shaw et al. 2008].

Simple scaling suggests that the risk of a YME repository completely filled with spent nuclear fuel from LIFE engines is slightly more than that of a repository completely filled with spent nuclear fuel from a fleet of light-water reactors, but still well within the statutory limits

established with 10 CFR 63, as shown in Figure 77. Scaled dose from a Yucca-Mountain-like repository containing LIFE waste is of the same order of magnitude as the Yucca Mountain TSPA-LA results, and well below the proposed regulatory limit. Major contributors to long term dose from LIFE waste are ¹²⁹I, ¹³⁵Cs, and ²⁴²Pu. We will need to do performance assessment calculations using accepted software tools (e.g., GoldSim) in the future [Shaw et al. 2008].

Other Alternatives

Another sub-critical system that has been proposed for the burning of spent nuclear fuel is known as accelerator transmutation of waste (ATW). In this process, reprocessing is used to separate TRU from LWR SNF for burning in a lead-bismuth cooled reactor. In contrast, SNF can be burned in LIFE without significant isotopic enrichment or chemical reprocessing. A side-by-side comparison of two proposed fuel cycles, one with ATW and the other with LIFE engines is shown in Figure 78 [Farmer 2008].

Safe Handling of Tritium

LIFE requires safe on-site tritiuin storage with solid-state tritide systems, which have been demonstrated on relatively large scale. The storage capacity of various solid-state hydrogen and hydrogen-isotope storage systems are compared in Figure 79 [DeLuchi 1989]. Solid-state tritium storage media must be selected that will release the tritium with relatively mild temperature swings near ambient temperature. The pressure-temperature characteristics of several candidate solid-state storage systems are compared in Figure 80. In Mg-Ni system may be an attractive system, with both high storage capacity, and modest operating temperature.

Cost and Availability of Advanced Materials

Cost of Advanced Steels

A cost model was developed and used to predict the cost to produce nickel-based alloys, including Type 316 stainless steel, as well as nickel-based Alloys C-276 and C-22. This cost model used raw materials data compiled by the United States Geological Survey (USGS), and represented graphically in the following Figure 81. The cost of raw materials required for the production of ferritic and austenitic steels, nickel-based alloys, and various refractory alloys. These costs were provided by the United States Geological Survey (USGS).

Requirements of Lithium & Beryllium for LIFE Reactor

Substantial quantities of beryllium and lithium will be required to build and operate LIFE plants. Typical inventories of beryllium and lithium for one selected LIFE engine design are: 14 to 15 MT of beryllium; and 6.24 to 7.75 MT of lithium. Typical inventories for the entire plant are slightly greater: 23.5 MT of beryllium; and 18.72 to 23.25 MT of lithium. The availability of

these strategic materials must be evaluated to ensure that there is enough material to warrant the continued development of LIFE technology.

Availability of Beryllium

Legislation and government programs that impact the availability of Beryllium are described by the Minerals Yearbook published by the United States Geological Survey. The Defense Production Act Title III provided for a \$9M Contract to Brush Engineering Materials (BEM) Incorporated, and enabled the construction of a new facility to make metallic beryllium products. The completion of construction is expected in the 2009-2010 timeframe. The National Defense Stockpile (NDS) was established so that the United States could maintain stockpiles of strategic materials such as beryllium for use during national emergencies. The NDS goal for December 31, 2006 was 45 metric tons of metallic beryllium powder, consolidated with hot isostatic pressing (HIP). This is in contrast to the 155 metric tons of beryllium metal proposed in the 2004 NDS Requirements Report to Congress in 2003. The USGS believes that the United States will remain self-sufficient in beryllium production [Shedd 2007].

BEM has reported proven bertrandite ore (Be₄Si₂O₇(OH)₂) reserves in Joab County, Utah of about 5.94 million dry metric tons with an average grade of 0.267% beryllium. This source represented about 15,900 metric tons of beryllium metal. If calculated more accurately, it is about 15,593 metric tons based upon the average grade.

During 1st Quarter 2007, BEM supplied Be blanks to the JET nuclear fusion reactor project. Other uses included metal-matrix composites for speakers, defense applications, tactical optics, including forward-looking IR systems, airborne electronics, space systems, and medical and industrial X-ray systems. Other exporters of Be include China and Kazakhstan.

The proven U.S. reserves, U.S. mine shipments, and world mine shipments of beryllium, expressed in metric tons (MT), are summarized in Figure 82. The corresponding number of LIFE engines that can be built from proven US reserves, US mine shipments, and world mine shipments of beryllium, are summarized in Figure 83. The proven reserves of 15,593 metric tons of beryllium metal in Joab County, Utah (USGS 2006) are sufficient to build approximately 664 LIFE plants, assuming that each system would require 24.5 metric tons of metallic beryllium. Approximately 155 metric tons of beryllium were shipped from mines within the United States in 2006, which was enough for the construction of approximately 7 LIFE plants. Annual worldwide mine shipments would be enough to build about 8 LIFE plants. The estimated 2006 US Be consumption of 91 MT Be Metal was valued at \$28 million. This corresponds to a cost of approximately \$308 per kilogram (\$139 per pound).

Availability of Lithium

Annualized 2006 lithium consumption in the US is estimated to be approximately 2,500 MT Li Metal, and was approximately 32% greater than the consumption in 2004. The single lithium

carbonate (Li₂CO₃) producer within the US appears to be Chemetall Foote Corp (US subsidiary of German Chemetall GmbH, which is owned by Rockwood Holdings of Princeton, NJ) [Ober 2007].

Chemetall Foote Corp produces lithium carbonate from brine wells near Silver Peak, Nevada. Chemetall Foote Corp also has other lithium operations in Silver Peak, Nevada; New Jacksonville, Tennessee, and Kings Mountain, North Carolina. Other countries involved in lithium production include: Canada, Chile, Argentina, Brazil, Portugal, China and Zimbabwe. Chile has been the world's leading producer of lithium carbonate since 1997, the year it first surpassed the US in production. Production in Chile was from two brine operations on the Salar de Atacama in the Andes Moutains. Concentrated brines were processed at two plants in Antofagasta. Chemetall Foote's subsidiary in Chile is Sociedad Chilena de Litio Ltda. and is involved in this production. Australia produces about 60% of the world's supply of solid lithium minerals at the Sons of Gwalia Ltd.'s Greenbushes spodumene mine. According to Austrialia's Department of Industry and Resources, this is the largest spodumene (LiAl(SiO₃)₂) deposit in the world. Canada's Tantalum Mining Corporation Ltd. (subsidiary of Hudson Bay Mining Company) also operates a substantial spodumene mine and concentrating plant near Bernic Lake in Manatoba

In regard to competing markets for lithium, lithium-ion and lithium polymer gel batteries appear to have the greatest potential (by large margin) for growth. The world market for these rechargeable batteries was \$4 billion in 2005, and is expected to grow to tens of billions (perhaps \$75 billion) in the next few decades. This market has been lost almost completely to the Chinese due to differences in labor cost between China and other countries, including both the US and Japan. The second largest use for lithium is the manufacture of rechargeable lithium batteries, due to the proliferation of wireless communications and computing. This may become the leading use in the near future.

Other well-known markets for lithium include lithium compounds used as catalysts and reagents for the production of synthetic rubbers, plastics and pharmaceuticals. For example, N-butyllithium is used to initiate the styrene-butadiene reaction necessary for the production of abrasion-resistant rubbers which require no vulcanization. Lithium hydroxide monohydrate is also a key ingredient in multipurpose greases.

The proven U.S. reserves, U.S. consumption and exports, deducting imports, and world production of lithium, expressed in metric tons (MT) are summarized in Figure 84. The number of LIFE engines that can be built from proven U.S. reserves, U.S. consumption and exports, deducting imports, and world production of lithium are summarized in Figure 85. Approximately 32,688 LIFE plants could be built with proven 2006 lithium reserves in the United States (750,000 MT). Approximately 32 LIFE plants could be built with 2006 domestic mine production (740 MT), while 903 LIFE plants could be built with 2006 worldwide shipments of lithium (21,000 MT).

Gaps in Knowledge & System Vulnerabilities

The irradiation levels in various parts of the LIFE engine have been predicted and are presented in Table 5. The TRISO fuel materials will receive approximately 240 dpa after 40 years, while the ODS steel structure will receive 210 dpa after only 6 years, which indicates periodic change-out of the steel chamber.

The experience base for TRISO fuels is limited, with 8 to 20% FIMA demonstrated with low enriched uranium (LEU), and 79% FIMA demonstrated with high enriched uranium (HEU) as shown in Figure 16. These LEU and HEU fuels experienced only approximately 3.5×10^{21} n/cm², but at a relatively high temperature of 1100 to 1200°C. In contrast, LIFE fuels will experience a much higher neutron dose of approximately 1.2×10^{23} n/cm², but at a much lower temperature of about 700°C. There is no operating experience with TRISO fuels at such high neutron fluences.

Little or no relevant experience exists at 1.2×10^{23} n/cm² with other possible high burn-up solid fuels, such as solid hollow-core (SHC) fuels and inert matrix fuels (IMF). Little experience exists with liquid fuels, with on-line reprocessing for the continuous removal of rare-earth fission products, which is necessary to prevent the precipitation of plutonium from the molten salt.

No known process exists for the re-configuration of LWR SNF into high burn-up TRISO, SHC or IMF for use in the LIFE engine. Fabrication processes for the fiber wrapped SHC fuel must be developed.

While predictive models have now been developed for structural, first wall and fuel materials, there is insufficient test data to calibrate model parameters, and validate model predictions. Data from irradiation, mechanical and corrosion tests are desperately needed.

Fundamental data on the reactive transport of fission products in various regions of the LIFE fuel and engine is required. The interaction of various fission products with encapsulation materials used in solid must be better understood.

There are a number of outstanding challenges for the first wall, the neutron multiplication blanket, the structural materials, and the high-temperature molten-salt coolants. Practical realization of the spherical ODS chamber required for construction of the LIFE engine also requires substantial work. For example, methods for the large-scale production of high quality ODS steel must be developed. Methods for making joints (welds) with high integrity are also required. Critical properties must be determined, and include: long-term creep rates and rupture times, fatigue, creep-fatigue and fatigue crack growth, compatibility and corrosion rates in molten salts, and environmental cracking. The effects of long-term exposure to high-flux fast neutrons at high temperature, the exact role of oxide nanoparticles in strengthening ODS steels, and the evolution of these small oxide particles during operation must all be better understood.

Strategy for Future Work

Over the next twelve months, the LIFE Materials Team will deliver more detailed predictive models for structural and first wall materials, as well as for solid and liquid fuels, a detailed plan for experiments necessary to calibrate and validate model parameters, and preliminary results from the initial materials performance experiments, assuming that adequate resources are available for the planned work scope. Specific issues that must be dealt with include: stresses associated with the accumulation of fission gases within the particles of fuel surrounding the fusion target area; pulsed heating of the fuel and first wall, and the associated mechanical stresses; high-temperature creep of structural materials; corrosion and environmental cracking of those reactor components exposed to the high-temperature molten-salt coolant and tritium breeding blanket; radiation damage on fuels, structural and first wall materials; the creation of appropriate test environments for materials within the laboratory, including extreme neutron fluxes with representative energy spectra, rapid thermal pulsing with homogeneous heating; exposures to high-temperature lithium-beryllium-fluoride molten salts, and special problems associated with the use of beryllium multipliers in such applications. Planning, modeling and early experimental work will be used to begin solving materials problems currently believed to challenge the LIFE reactor concept. A detailed research and development strategy will be formulated, thereby enabling the team to attack crucial materials problems in an efficient and comprehensive manner. The initial fabrication of prototypical fuels by an experienced industrial partner will be initiated, incorporating those design changes necessary to allow the fuels to survive higher burn-up and neutron dose. The proposed developmental work will leverage the extensive knowledge base within the Department of Energy that is now being Advanced Gas Reactor (AGR), and will provide impetus for creating fuels capable of even higher burn-up, and capable of accommodating a much broader range of uranium feeds.

This section of the Topical Report (TR) describes work that will be done to address the materials challenges that have been identified and discussed. Materials research and development activities will focus on: fuel design, fabrication, and performance; and structural and first wall and structural materials options, fabrication and performance; specialty materials for the balance of plant; materials availability and cost; and issues pertaining to the end of the fuel cycle, including proliferation resistance, interim storage, and geological disposal of the residual waste.

A systematic approach to materials modeling will enable design of next-generation materials, as illustrated in Figure 76. Accelerated testing of materials will be required, including irradiations in a new three-beam accelerator facility at LLNL, as illustrated in Figure 77

The LIFE Engine requires materials that can withstand harsh conditions of radiation, high temperature, active chemical agents, and pulsed (20/s thermal and mechanical stress for periods of time as long as 60 years. While we have identified materials and materials systems candidates for first wall and fuel, practical experience and modeling code validation regarding these

materials under the extreme conditions of LIFE does not yet exist. Even if an appropriate neutron material testing facility were to exist, testing materials over the intervals of 60 years will never be a viable option. Accelerated testing methods coupled to advanced theory and modeling is the only way to rapidly evaluate candidate materials. Thus, development of capabilities for reliable accelerated material testing becomes a critical priority for the development of LIFE.

As part of the point design effort we will develop detailed plans for a multi-ion-beam accelerated materials radiation facility while carrying out early experimental simulations of reactor environment radiations by sequential materials irradiations with helium and heavy ions to simulate the materials property changes associated with the LIFE environment. Four key materials will be studied: tungsten, which will be used to fabricate the solid first wall; oxide dispersion strengthened ferritic steel, the structural material for engine construction; beryllium, the proposed neutron multiplier; and graphite, the proposed as a reflector in the baseline engine design. The ODS steel will be studied to radiation damage levels in excess of 200 dpa. Silicon carbide will be studied at radiation damage levels representative of pure fissile and pure fertile fuels. Temperatures for these irradiations will be representative of the anticipated operating temperatures for LIFE.

Integral with this experimental simulation will be the implementation of existing and the development of advanced modeling codes to interpret the experimental data. More importantly the successful implementation of the modeling codes will also serve as a confidence builder for prediction of material performance under LIFE conditions.

The degradation and failure of LIFE Engine components will be limited by high-temperature corrosion in Li₂BeF₄, LiNaBeF₄ and other coolants. A testing program will be needed to understand the corrosion, environmental fracture, thermally-driven fatigue, and radiation damage of hybrid reactor materials in high-temperature molten salts with high neutron fluence. Corrosion and environmental cracking will be exacerbated by thermal aging and radiation damage of crystalline solids, and the formation of transmutation products such as tritium fluoride and related chemical species (TF, DF & HF). Galvanic coupling of dissimilar materials, with one serving as a catalytic cathode, will drive anodic dissolution of the other at an accelerated rate. Enhanced mass transport in the intense flow-field of the vortex and annuli will accelerate corrosive dissolution of materials.

Fluoride salts formed with Mo, W, Ni and V are less stable than TF (HF) which will be formed by transmutation of lithium. Therefore, these metals are believed to have good corrosion resistance in FLIBE. In contrast, the other metals would be expected to undergo reaction with the fluoride ion to form stable fluoride-salt corrosion products.

Materials have been selected for construction of a LIFE Engine, and the susceptibility of those materials to corrosion must be determined. When problems are identified, alternative materials designs and selections may be required.

These tasks include: nuclear burn and neutron activation, fluence, and dpa analyses; assessment of the TRISO fuel designs with determination of their relative abilities to reach extended burnups, specifically TRISO pebbles with WG-Pu or HEU for 5-10 year LIFE engine operation; quantitative assessment of the performance and lifetime of structural and first-wall materials; and a preliminary cost analysis of the system

As part of the point design effort we will develop detailed plans for a multi-ion-beam accelerated materials radiation facility while carrying out early experimental simulations of reactor environment radiations by sequential materials irradiations with helium and heavy ions to simulate the materials property changes associated with the LIFE environment.

The design of a long-lived machine of any type capable of complete burn-up will require designs based upon predictive models that must be validated and calibrated. Even if an appropriate neutron material testing facility were to exist, testing materials over the intervals of a decade or more will never be a viable option. Accelerated testing methods coupled to advanced theory and modeling is the only way to rapidly evaluate candidate materials. Thus, development of capabilities for reliable accelerated material testing becomes a critical priority for the development of LIFE. Integral with this experimental simulation will be the implementation of existing and the development of advanced modeling codes to interpret the experimental data...

Service Life Prediction Strategy

- 1) Documentation of the irradiation testing strategy for prototype fuel particles that will be used in FY09 and beyond, including a detailed program to integrate multiple pre-exiting particle accelerators to simultaneously damage materials and implant species such as helium.
- 2) Collect published reports and articles on materials required for solid fuel fabrication, including but not limited to: fertile and fissile materials, in both metallic and ceramic forms; silicon carbide; zirconium carbide; carbons and graphite; and graphite fibers with refractory coatings such as silicon carbide; and molten-salt resistant cladding materials
 - a) Establish database from published data
 - b) Use PARFUME code to predict TRISO performance
 - c) Obtain other established models to predict fuel performance
- 3) Develop detailed models for high burn-up solid fuels (TRISO, SHC & IMF) accounting for:
 - a) Stresses due to fission gas accumulation and thermal pulsing
 - b) Transmutation and the effects of such compositional change on properties
 - c) Irradiation-induced structural damage on fuel performance
 - d) Chemical reactivity of materials with high-temperature molten salts
- 4) Apply those models/codes to predict the failure mechanism of solid fuels under assumed conditions representative of the conceptual LIFE engine
 - a) Weapons-Grade Plutonium (WG Pu)
 - b) High Enriched Uranium (HEU)
 - c) Natural and Depleted U (DU)
- 5) Identify predictive modeling needs
 - a) Specific data required as inputs to codes
 - b) Advanced physics methodologies for predicting the evolution of materials under extreme environments, and over prolonged periods of time, accounting for the stochastic-nature of the problem, and leveraging advances in supercomputing
- 6) Develop thermochemical model of liquid fuel
 - a) Predict the solubilities of uranium and plutonium
 - b) Predict the solubilities of rare-earth (RE) and noble-metal (NM) fission products
- 7) Comprehensive suite of models for prediction of anticipated failure modes in solid fuels
 - a) Calibrated model parameters for the various models
 - b) Documented model validation with existing experimental database
- 8) Predictions
 - a) Time-dependent temperature distribution in solid fuels
 - b) Lifetime of solid fuel designs under expected LIFE conditions
- 9) Sensitivity analysis determining the relative effects of temperature, radiation, fission gas buildup, and molten-salt corrosion on fuel lifetime of solid fuels

- 10) Assessment of solid fuel designs with determination of their relative abilities to reach extended burn-ups and withstand high damage-rates, as well as helium and hydrogen production in LIFE operating conditions
 - a) Identification of suitable liquid fuel compositions to meet LIFE missions
 - b) Conceptual design for continuous on-line process for removal of RE/NM fission products
- 11) Develop credible models for predicting the performance of structural materials:
 - a) Thermal pulsing and associated stresses
 - b) Corrosive attack by the high-temperature molten salts and transmutation products
 - c) Thermal creep
 - d) Radiation damage
- 1) Credible conceptual mechanistic model for:
 - a) Thermal pulsing and associated stresses
 - b) Corrosive attack by molten salts
 - c) Thermal creep
 - d) Radiation damage
- 2) Quantitative predictive model for:
 - a) Thermal pulsing and associated stresses
 - b) Corrosive attack by molten salts
 - c) Thermal creep
 - d) Radiation damage
- 3) Library of calibrated model parameters:
 - a) Thermal pulse model
 - b) Corrosive model
 - c) Thermal creep model
 - d) Radiation damage model

Accelerated Testing Strategy

- 1) Develop a detailed materials development and test plans
 - a) Experimental work to address anticipated degradation modes in materials
 - b) Determination of the effects of pulsed surface heating
 - i) Laser heating experiment
 - ii) Pulsed current experiment
 - c) Determination of the effects of radiation damage and swelling
 - i) Irradiations with single-beam accelerators
 - ii) Irradiations with triple-beam accelerator
 - iii) Collaborative testing in fast-flux reactors
 - d) Long-range irradiation tests in the LIFE Demonstration Unit
- 2) Necessary database to support to support development of conceptual and predictive models
 - a) Thermal pulse model
 - e) Corrosive model
 - f) Thermal creep model
 - g) Radiation damage model
- 4) Procure or synthesize structural materials for testing
 - a) First wall materials: tungsten, tungsten carbide
 - b) Structural materials: ODS FM steels, TMT steels, amorphous metals
 - c) Corrosion resistant coatings: tungsten, vanadium, amorphous metals
 - d) Neutron multipliers: beryllium, beryllium oxide
 - e) Coolants: Li-Na-K-F. Li₂BeF₄ and Li-Pb
- 5) Experimentation with molten salt and liquid metal coolants
 - a) Determination of credible coolant properties as a function of temperature
 - b) Determination of credible coolant properties as a function of composition
 - c) Predictive equations for properties as a function of temperature and composition
- 6) Experimentation with structural materials
 - a) Perform corrosion and environmental cracking tests
 - b) Perform thermal-pulse simulation experiment (TPSE)
 - c) Measure the hardness and strength of irradiated samples
 - i) Ion beam irradiation at high fluence with particle accelerator
 - ii) Initial neutron irradiations at relatively low fluence with research reactor
- 7) Radiation effects testing for structural and first wall materials
 - a) Initial accelerated testing with triple-beam accelerator
 - b) High-fluence neutron effects in reactor
 - c) High-fluence neutron effects LIFE demonstration unit
 - d) Determine the feasibility of performing neutron irradiations at external facilities
- 8) Data for further calibration and validation of structural material models

- a) Pulsed heating experiment with laser
- b) Pulsed heating experiment with current
- c) Coolant properties as function of temperature
- d) Corrosion rates as function of temperature and radiation damage
- e) Hardness, strength and creep as function of temperature and radiation damage
- f) Ion beam irradiation at high fluence with particle accelerator
- g) Initial neutron irradiations at relatively low fluence with research reactor
- 9) Radiation effects testing for solid fuel materials
 - e) Initial accelerated testing with triple-beam accelerator
 - f) High-fluence neutron effects in reactor
 - g) High-fluence neutron effects LIFE demonstration unit
 - h) Determine the feasibility of performing neutron irradiations at external facilities
- 10) Quantitative assessment of the performance and life time of structural and first-wall materials
- 11) Fabrication of advanced prototypical fuels for testing
 - a) Enhanced TRISO fuels in collaboration with BWXT, ORNL and INL
 - b) New SHC fuels in collaboration with additional partners experience in composites
 - c) Inert Matrix Fuels in Collaboration with CEA
- 12) Develop an on-line process for continuous removal of RE and NM fission products
- 13) Testing of high burn-up solid and liquid fuels in LIFE demonstration unit

Collaborators

In addition to the work to be performed at LLNL, we also propose to take advantage of the unique expertise available at other national laboratories. The work and the contracts described below are included in the Task list description below.

BWXT

They would participate and contribute in the fuels fabrication area, and would provide a conceptual process design for manufacture of the fuels based on WG Pu and HEU. They are the only facility in the United States with TRISO production capability, and qualified to handle HEU. We propose a contract of \$50K.

UC Berkeley

We are already drawing upon their experience in the safety analysis, and want to expand that effort to include nuclear fuels and radiation damage. We propose a contract of \$75K.

Caltech

CalTech will work on the development of multi-scale models of Pu and HEU fuel performance for 99% FIMA and thermo-mechanical stresses associated with the pulsed operation of LIFE fuel pellets. We propose a contract \$75K.

Summary

In summary, we have discussed a concept for a hybrid fusion-fission energy engine. Significant challenges in demonstrating the physics of IDFI, development of advanced materials and DPSSL technologies remain. However, recent progress on NIF at LLNL and on the physics and technologies of inertial confinement fusion, DPSSL technologies and materials science for applications in the nuclear environment in the US and elsewhere around the world promise to bring us closer to realizing the longstanding vision of fusion-fission hybrids. Starting with a laser fusion driver with gain of ~ 50 , such a device would be capable of burning large quantities of spent nuclear fuel, natural or depleted uranium or weapons-grade Pu to generate GW of thermal power in a subcritical fission blanket. It would therefore provide pathways to energy generation in a closed nuclear fuel cycle and to eliminating the excess inventory of spent nuclear fuel on the way to pure fusion energy.

LIFE can burn excess plutonium in a high-gain blanket, and is being explored for NA-22. This conceptual system is assumed to be fueled with 7 tons of weapons-grade plutonium (WG-Pu). It is further assumed that: TRISO kernels are blended 80% ZrC + 20% Pu; TRISOs are loaded with 400 wppm boron as burnable poison; fusion power is 375 MW (37.5 MJ at 10 Hz); the flat-top thermal power is 3000 MW; and the blanket gain is \sim 8. When fueled with weapons-grade Pu or HEU, LIFE will typically be fueled with \sim 6 or 7 metric tons and with \sim 350-400 MW fusion provide 3000 MWth or 1350 MWe for about 6 years while burning the Pu to over 99% FIMA.

An early LIFE demonstration unit, burning WG-Pu, will enable the development of an advanced power system capable of helping provide much of the Nation's future energy needs, with substantial reductions in the generation of high-level radioactive waste. Specific system improvements that are anticipated include the advent of fast-ignition ICF, advanced solid and molten salt fission blankets, and advanced structural materials capable of operating at higher temperatures. The WG-Pu mission has the potential of enabling the hybrid LIFE systems, and will also enable pure fusion, since LIFE is a practical and realizable precursor for pure fusion.

The design of LIFE engines capable of meeting the identified missions will require data and models that account for the corrosion and environmental cracking of as-received and irradiated structural and cladding materials in high-temperature molten salt coolants and liquid fuels.

Several topics and issues related to the development of high burn-up fuels for the sub-critical fission blanket in a LIFE Engine have been addressed in this quarterly report and include: detailed description of the structure of TRISO, SHC and IMF fuels, and the processes required for their fabrication; radiation swelling of silicon carbide and graphite used in the fuels; static and dynamic stresses associated with the accumulation of fission gas and thermal pulsing; stress analysis with ALE3D and ABAQUS finite element models; thermal migration of the uranium oxycarbide kernel; attack of the silicon carbide capsule with fission products such as palladium; and the abrasion and corrosive attack of clad pebbles. Methods for mitigating identified problems

include: operation of the LIFE engine at a lower operating temperature to avoid the non-saturable void-swelling regime; designing the TRISO fuel with a thicker SiC capsule; using ZrC as a getter to capture liberated oxygen; substituting UOC for UO_2 to prevent kernel migration; and using sacrificial SiC is used close to the kernel to capture threatening fission products such as palladium.

Several topics have also been addressed related to the structural and first wall materials. New oxide dispersion strengthened (ODS) ferritic-martensitic (FM) steels such as 9Cr-ODS & Kobe 12YWT have exceptional tensile strength at high temperature, have been used successfully use for cladding fuel pins in the BOR-60 sodium-cooled fast reactor ODS has been successfully joined using several different industrial welding processes, and provides exceptional high-temperature properties LIFE engine design. Based upon published corrosion rates for other chromium containing steels, a reasonable estimate for the corrosion rate of ODS by molten fluoride salts believed to be on the order of 1 mil per year.

The LIFE engine will produce substantially less HLW and weapons-attractive material (Pu and TRU) than an LWR of comparable capacity. It appears that an interim storage facility and deep geological repository can be built for LIFE SNF with adequate thermal management, capable of preventing waste packages from exceeding the allowable operating temperature for container materials. Tritium inventories can be managed with well-known and characterized solid state storage media. There are adequate supplies of Li and Be to build and supply the required fleet of LIFE engines.

The LIFE Engine is an advanced high-performance nuclear energy conversion device with an ICF spark plug and requires integration of some of the world's most advanced reactor materials. The initial engine concept includes a W 1st wall, ODS steel structure, Be neutron multiplier blanket, TRISO fission blanket, and molten salt coolant.

There are three primary missions for the LIFE engine, which include the burning of excess weapons-grade plutonium (WG-Pu), natural and depleted uranium (DU and HEU), and lightwater reactor (LWR) spent nuclear fuel (SNF). Materials in the LIFE Engine will be subjected to a severe radiation and high temperature environment. Such operating conditions will challenge current materials and stimulate development of next-generation materials. The LIFE engine's structural challenges include: need for high-temperature strength; resistance to high-temperature creep; immunity to radiation damage, including swelling and helium embrittlement; resistance to corrosion and environmental cracking in high-temperature molten fluoride salts; and the ability to be fabricated into necessary shapes and configurations with practical welding processes. It is hoped that these challenges can be met with oxide dispersion strengthened (ODS) ferritic steels, sheets and coatings of refractory metals such as tungsten, advanced solid or liquid fuels, and other advanced reactor materials.

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Tables

Table 1 – Budget for Proposed Statement of Work

Task No.	Task	Funding Request \$K	Procuremen ts FY08 \$K	Task Total \$K
1	LIFE proliferation resistance, enterprise cost model and technology comparisons	900		900
2	LIFE engine options and performance optimization	1400	200	1600
2.1	Neutronics and engine design and contract to CalTech	600	75	675
2.2	Fuel fabrication and performance, and contracts to University of California at Berkeley, BWXT	600	125	725
2.3	First wall and structural materials options	200		200
	Total	2300	200	2500

Table 2 – Composition of 40 Tons of DU Fuel in LIFE Engine at 90 to 99.9% FIMA

	Burn-up			
Isotope	90%	95%	99%	99.9%
235U	1.6 kg	120 g	560 mg	36 mg
238U	3300 kg	1800 kg	490 kg	150 kg
237Np	5.4 kg	1.9 kg	290 g	69 g
239Pu	210 kg	58 kg	11 kg	3.2 kg
241Pu	10 kg	21 kg	3.6 kg	1.1 kg
241Am	4.2 kg	270 g	13 g	1.9 g
246Cm	150 kg	170 kg	120 kg	79 kg

Table 3 – Composition of 7 Tons of WG-Pu Fuel in LIFE Engine After Burn

Isotope	Initial mass	Final mass
239Pu	6.56 tons	1.3 mg
240Pu	406 kg	<1 mg
241Pu	9.1 kg	1.2 mg
242Pu	1.4 kg	<1 mg
241Am	15.4 kg	<1 mg
Total actinides	6.99 tons	2.83 kg (2.04 kg 246Cm)

Table 4 – Performance Characteristics of Various Types of LIFE Plants

			1.5	1.0	2.4			
		Daga	1.5 GWe	1.0 GWe	GWe	DU +	Pu	Dura
Crystam Chamatanistia	I Inita	Base	NIF FI		HS 2ω			Pure IFE
System Characteristic	Units Hz	Case 13.33		LIA FI		IFE T ₂	Burner	
Repitition Rate	W	2	13.33	15.00	10.00	10.00	13.33	15.00
C-Laser Frequency				1.09			2	1.12
C-Laser Energy	MJ	0.659	0.809		2.702	0.659	0.659	
C-Laser Pulse Width	ns	25	25	25	25	25	25	25
I-Laser Frequency	ω	1	1	1	1	1	1	1
I-Laser Energy	kJ	150	150	150	0	150	150	150
I-Laser Pulse Width	ps	10	10	10	10	10	10	10
Chamber Radius	M	2.25	3.03	2.44	4.50	2.90	2.39	4.50
TRISO Pebble Dia.	cm	2.00	2.00	2.00	2.00	2.00	2.00	2.00
TRISO Particle Dia.	mm	1.00	1.00	1.00	1.00	1.00	1.00	1.00
TRISO Kernel Dia.	Mm	0.60	0.60	0.60	0.60	0.60	0.60	0.60
Be Pebble Dia.	cm	2.00	2.00	2.00	2.00	2.00	2.00	2.00
Fusion Target Yield	MJ	37.5	68	44	150	37.5	37.5	150
Fusion Target Yield	n/cm ² /s	1×10 ¹⁵						
Fusion Power	MW	500	907	660	1500	375	500	2250
Fuel Composition		DU	DU	DU	DU	DU	Pu	T
Fission Fuel	MTHM	40	73	47	160	67	7	0
Thermal Power	MW	2000	3627	2640	6000	2500	3000	2655
Burn-up	%FIMA	99.5	99.5	99.5	99.5	99.5	99.5	99.5
Maximum k-effective	k _{eff}	0.9	0.9	0.9	0.9	0.9	0.9	0.9
Service Life	years	50.00	50.00	44.44	66.67	66.67	5.00	50.00
Availability	fraction	0.9	0.9	0.9	0.9	0.9	0.9	0.9
Coolant Composition		FLIBE						
Coolant Inventory	MT	40	40	40	40	40	40	40
Fusion Gain	G_{fusion}	57	84	41	56	57	57	134
Fission Gain	$G_{ m fission}$	4	4	4	4	6.667	6	1
Total Gain	G _{total}	228	336	162	222	379	341	134
DPSSL Efficiency	%	10%	10%	10	10	10	10	10
Thermal to Electric	%	45	45	45	45	45	45	45
Re-circulating Power	%	9.76	6.61	13.72	10.01	5.86	6.51	14.1
Gross Electric Power	MWe	900	1632	1188	2700	1125	1350	1195
Auxiliaries	MWe	25	25	25	25	25	25	25
Net Electric Output	MWe	787	1499	1000	2405	1034	1237	1001

Note: C-Laser = Compression Laser; I-Laser = Ignition Laser.

Table 5 – Fluence and Damage Levels in the LIFE Engine

	First Wall		Be Blanket	TRISC	Fuel	
	Tungsten	ODS Steel	Beryllium	SiC	Graphite	
Fast Fluence (n/cm²)	2.89E+22	2.89E+22	2.28E+22	7.66E+20	7.66E+20	
Total Fluence (n/cm²)	8.00E+22	8.00E+22	7.87E+22	2.97E+21	2.97E+21	
Damage (dpa)	10	36	13	6	3	
He (appm)	10	312	5000	88		
H (appm)	22	1176	51	139		
Temperature (°C)	750-1300	650-750	650-750	750-800	750-800	

Note: The irradiation levels in various parts of the LIFE engine have been predicted and are presented in Table 1. The TRISO fuel materials will receive approximately 240 dpa after 40 years, while the ODS steel structure will receive 210 dpa after only 6 years, which indicates periodic change-out of the steel chamber.

Table 6 – Density and Melting Points of Candidate Fissionable Fuels

Uranium	Density (g/cm ³)	Melting Point (°C)	Analog	Structure
Compound				
UN	14.32	2850	NaCl	Cubic
UO_2	10.97	2880	CaF ₂	Cubic
UC ₂	11.68	2470		Tetragonal
UC	13.63	2370	NaCl	Cubic
USi	10.40	1600		Orthorhombic
UAl ₂	8.10	1590		
UAl ₃	6.70	1320		
U	19.13	1133		
Th ₄	6.32	1114		
UF ₄	6.70	1036		
UAl ₄	6.00	730		

Note: A wide variety of potential fuel materials exist for solid and liquid fuels. In the case of solid fuels, nitrides, oxides, carbides, silicides, aluminides and metals have all been considered. As shown in Table 2, oxides, nitrides and carbides have the highest melting points. Commercial TRISO fuels use a mixed ceramic phase, uranium oxy-carbide (UOC), which minimized thermal migration effects of the kernel.

Table 7 – Challenges Facing TRISO Fuels & Mitigation Strategies

Challenges facing TRISO fuels	Design feature meeting the challenge
Irradiation swelling of SiC capsule & PyC layers	The LIFE engine is designed with a lower operating temperature to avoid the non-saturable void-swelling regime
Stress in SiC capsule due to fission gas (Kr & Xe) pressure	The SiC capsule is designed with a thicker wall
Reaction of C & O ₂ released from UOC to form CO	ZrC is used as a getter is used to capture liberated oxygen
Fuel kernel migration up temperature gradient	UOC is substituted for UO ₂ in fuel kernel
Fission product (FP) attack of SiC capsule	Sacrificial SiC is used close to kernel to form stable 1:3:3:5 U:Pd:Si:C (MP ~ 1952°C) Use of diffusion barrier to FP prevent contact

Note: Several challenges must be met to achieve high burn-up with TRISO-type fuels in the LIFE engines. The challenges that must be met, and known strategies for meeting these challenges are outlined in Table 3.

Table 8 – Blanket and Coolant Properties

Salt Formula	Formula Weight	Melting Point	Latent Heat of Fusion	Vapor Pressure (900°C)	Density	Heat Capacity (700°C)	Viscosity	Thermal Conductivity	Neutron Capture vs. Graphite	Moderating Ratio
	FW	MP	ΔH_{f}	P_{vap}	P	$\rho \times C_p$	μ	K		
	(g/mol)	(°C)	(cal/g)	(mm Hg)	(g/cm ³)	(cal/cm ³ /°C)	(cP)	(W/m-K)		
Li	6.9	179			554					
Be	9.0	1283			1820					
Na	23.0	97.81			970					
K	39.1	63.65			860					
LiF	25.9	842			2300					
BeF ₂	47.0				1086					
LiF-BeF ₂	33.0	460		1.2	1.94	1.12	5.6	1.00	8	60
LiF-NaF-BeF ₂	38.9	315		1.7	2.00	0.98	5.0	0.97	20	22
KF-ZrF ₄	103.9	390		2.8	2.80	0.70	5.1	0.45	67	3
Rb-ZrF ₄	132.9	410		1.3	3.22	0.64	5.1	0.39	14	13
LiF-NaF-ZrF ₄	84.2	436		5 2	2.79	0.84	6.9	0.53	20	13
LiF-NaF-KF	41.3	454		0.7	2.02	0.91	2.9	0.92	90	2
LiF-NaF-RbF	67.7	435		0.8	2.69	0.63	2.6	0.62	20	8
72LiF-16BeF ₂ -12ThF ₄			63.00	0.1 torr	3.35		12	1.10		
70LiF-12BeF ₂ -18ThF ₄					3.87					
71LiF-2BeF ₂ -27ThF ₄			54.00		4.52		15-25	0.70		

Note: A wide variety of high temperature fluids exist that can be used to cool fission, fusion and fusion-fission hybrid systems, and include liquid metals and alloys, and fluoride salts. These include lithium, beryllium, sodium, potassium, alkali metal alloys, lithium fluoride, beryllium fluoride, lithium-beryllium fluoride (FLIBE = Li_2BeF_4), and mixed alkali-metal fluorides (FLiNaK = Li-Na-K-F). The presence of the lithium in the salt enables breeding of tritium, while the beryllium helps moderate neutrons. At the present time, the LIFE engine design specifies FLIBE for the primary coolant system, with the extraction of breed tritium from this fluid. The secondary cooling loop will use beryllium-free FLiNaK, which is both less expensive and less hazardous than the FLiNaK. The properties of many of these high-temperature heat-transfer fluids are summarized in Table 5 [Williams et al. 2006; Moir et al. 1985; Wright and Foster 1973; Rosenthal et al. 1971].

Table 9 – Important Physical Properties of Candidate Structural Materials

Property	Type 316L	Type HT-9	V-Cr-Ti
	Stainless Steel	Ferritic Steel	Alloy
Melting Temperature (°C)	1400	1420	1890
Density (g/cm ³)	8.0	7.8	6.1
Poisson's Ratio	0.27	0.27	0.36
Modulus of Elasticity (GPa at 400°C)	168	180	120
Linear Thermal Expansion (10 ⁻⁶ /K)			
400°C	17.6	11.8	10.2
500°C	18.0	12.3	10.3
600°C	18.3	12.6	10.5
Thermal Conductivity (W/m-K			
400°C	19.5	26.8	33.6
500°C	21.0	27.3	34.5
600°C	22.5	27.7	35.3
Electrical Resistivity (µm)			
400°C	1.01	0.91*	0.67
500°C	1.06	0.99*	0.74
600°C	1.12	1.05*	
Specific Heat (J/kg-K)			
400°C	560	600	535
500°C	575	680	560
600°C	580	800	575

Note: A number of steels have been considered for building the structure of the LIFE engine. These include a variety of austenitic stainless steels, ferritic steels, and refractory metal alloys such as V-Cr-Ti, which are all included in Table 4. The austenitic steels are prone to extreme swelling and high temperature thermal creep, as will be discussed subsequently. Ferritic steels show relatively little swelling during neutron irradiation, and can be enhanced by inclusion of a nano-dispersion of oxide particles. The refractory may be useful at higher temperatures than the levels that can be attained with austenitic and ferritic steels. The data for electrical resistance of the ferritic steel is assumed to be equal to that of Type 410 stainless steel (*) [Smith et al. 2000].

Table 10 – Nuclear Related Properties of Candidate Structural Materials

Property	Type 316L Austenitic Steel	Type HT-9 Ferritic Steel	V-5Cr-5Ti Vanadium Alloy	Silicon Carbide
Melting Temperature (°C)	1400	1420	1880	2600ª
Radiation Damage Rate (dpa)	11	11	11	12
Helium Transmutation Rate (appm)	174	130	57	1500
Hydrogen Transmutation Rate (appm)	602	505	240	560
Nuclear Heating Rate (W/cm ³)	11	11	7	11

Note: Austenitic 316L stainless steel, high-strength HT-9 ferritic steel, refractory V-Cr-Ti alloy, and silicon-carbide, all of potential interest for application in the LIFE engine have been irradiated, with values of the radiation damage rate, helium transmutation rate, hydrogen transmutation rate, and nuclear heating rate measured. Irradiations were done with ~ 14 MeV neutrons at a superficial power density of ~ 1 MW/m² for 1 year. These published data are tabulated in Table 5 [Smith et al. 2000].

Table 11 – Radiation Swelling of Inconel 600

Material	Specimen Fluence (n cm ⁻²)	Vol. Change by	Vol. Change by	Data
		Immersion (%)	TEM (%)	Source
Inconel 600	$1 \times 10^{22} \text{ n cm}^{-2} \text{ at E} > 0.1 \text{ MeV}$	3.6 %	5.0 %	Table 1
Inconel 600	$1 \times 10^{22} \text{ n cm}^{-2} \text{ at E} > 0.1 \text{ MeV}$	0.32%	< 0.1%	Table 1
Inconel 600	$1 \times 10^{22} \text{ n cm}^{-2} \text{ at E} > 0.1 \text{ MeV}$	< 0.5%		Fig. 11
Inconel 600	$4 \times 10^{22} \text{ n cm}^{-2} \text{ at E} > 0.1 \text{ MeV}$	1%		Fig. 11
Inconel 600	$7 \times 10^{22} \text{ n cm}^{-2} \text{ at E} > 0.1 \text{ MeV}$	2%		Fig. 11
Inconel 600	$9 \times 10^{22} \text{ n cm}^{-2} \text{ at E} > 0.1 \text{ MeV}$	5%		Fig. 11

Note: Data for the irradiation-induced swelling of Inconel 600, and nickel-based austenitic steel has been published by Makenas et al. more than 20 years ago, as shown in Table 6, and exceeds 5% at 9×10^{22} n cm⁻² and E > 0.1 MeV. Such swelling makes the nickel-based austenitic alloys is believed to make this family of alloys unacceptable for application in the LIFE engine. [Makenas et al. 1986, Table 1, Figure 11].

Table 12 – Guidance on Acceptable Compositions for Oxidation-Resistant Refractory Steels

Max. Temp. (°C)	Carbon (%)	Silicon (%)	Chromium (%)	Nickel (%)
850	0.35-0.45	2-3	8-10	
050	0.55 0.45	2 3	0 10	
900	< 0.12	< 0.8	16-18	
950	0.25-0.34	2-3	12-14	6-7.5
750	0.25 0.51	2 3	12 11	0 7.5
1000-1050	< 0.2	2-3	19-22	12-15
1100-1150	< 0.15	< 1	27-30	
1100 1130	0.15	- 1	27 30	

Note: The steels used for construction of the LIFE engine must be able to withstand high temperatures without melting or excessive thermal creep. General guidelines for the compositions of oxidation-resistant refractory steels have been published, as shown in Table 7. At temperatures between 850°C, which are the current upper bounds for the LIFE engine design, an alloy, with recommended concentrations of 0.35 to 0.45% for carbon, 2 to 3% for silicon, and 8 to 10% for chromium. The oxide dispersion strengthened (ODS) ferritic steels that are now being considered for the LIFE engine have comparable levels of chromium.

Table 13 – Materials Compatibility with Li₂BeF₄ Coolant

Alloy	Expected Corrosive Attack
V Alloys	No – Acceptable
W Alloys	No – Acceptable
Mo Alloys	No – Acceptable
Austenitic Steel	Yes – Manageable with Careful Engineering
Ferritic Steels	Yes – Manageable with Careful Engineering
Nb Alloys	Yes – Unacceptable
Ta Alloys	Yes – Unacceptable
Be Alloys	Yes – Requires Mitigation
Graphite	Unknown

Note: The corrosive attack of ferritic steels in FLIBE is considered to be manageable, provide that steps are taken to control the dissolution of the chromium into the fluoride salt. Corrosion resistance can be enhanced through careful engineering design, avoiding galvanic coupling of dissimilar metals, unless sacrificial protection is sought. For example, either Li or Be could be used as sacrificial anodes for ferritic steel components. The beryllium neutron multiplier should be protected with a cladding or encapsulation to prevent unacceptable rates of dissolution during the service of the LIFE engine.

Table 14- Free Energies of Formation for Various Fluorides

Fluoride Salts:	Free Energy of Formation
Possible Corrosion Products	(kcal/g-mol F) at 1000K
М.Б	50.2
MoF ₆	-50.2
WF ₆	-56.8
NiF ₂	-55.3
VF ₅	-58
VF ₄	-66
HF	-66.2
FeF ₂	-66.5
CrF ₂	-75.2
BeF ₂	-106.9
LiF	-125.2

Note: The transmutation of lithium in the Li₂BeF₄ forms corrosive tritium fluoride, which behaves chemically like hydrogen fluoride, or hydrofluoric acid. The ultimate selection of materials for long-term containment such corrosive transmutation products in fluid pumping loops requires the use of alloy systems with constituents that form fluoride salts less thermodynamically stable than hydrogen fluoride. It is therefore useful to rank candidate alloying constituents based upon the free energies of formation of their respective fluoride salts, each with the units of kilocalories per gram mole of fluoride at 1000 Kelvin. Such a ranking reveals that MoF₆, WF₆, NiF₂, VF₅, VF₄ have lower (negative) free energies than that of HF, indicating acceptability of Mo, W, Ni, and V, while FeF₂ has a (negative) free energy comparable to HF, indicating marginal acceptability of Fe-based alloys. Chromium has higher (negative) free energy, which explains the dissolution of chromium by such fluoride salts, and the formation of the well-known chromium depletion layer. Some refractory materials such as Nb-based and Ta-based alloys would be expected to be unacceptable for long-term exposure in high-temperature fluoride salts. Note that metallic beryllium (neutron multiplier in LIFE engine) would be expected to undergo dissolution in Li2BeF4, especially with galvanic coupling to more noble materials such as Mo-, W-, Ni-, V- and Fe-based alloys [Cheng et al. 2003].

Table 15 – Corrosion of Steels in High Temperature Molten Salts

Toop #	Alloy	Salt	Duration (hours)	T _{max} (°C)	Corrosion Depth (mil)	Corrosion Rate (mil/yr)	ONL Report [Reference]
116	316SS	FLiNaK	500	815	4.0	70.1	1294
119	316SS	FLiNaK + NaK	500	815	2.0	5.8	1294
347	Inconel	50NaF-50ZrF ₄	3000	815	11.0	32.1	1692
518	Inconel	NaF-ZrF ₄	3000	815	11.0	48.2	2338
346	Inconel	50NaF-50ZrF ₄	2000	815	9.0	39.4	1692
519	Inconel	NaF-ZrF ₄	2000	815	12.5	109.5	1692
78	Inconel	FLiNaK	1000	815	13.0	113.9	1294, 2337
	Inconel	NaF-ZrF ₄	1000	815	3.0	26.3	[76]
278	Inconel	NaF-ZrF ₄	1000	815	5.0	43.8	2338
399	Inconel	NaF-ZrF ₄	1000	815	10.0	87.6	2338
	Inconel	60NaF-40ZrF ₄	1000	815	5.0	43.8	CF-57-9-35
	Inconel	50NaF-50BeF ₂	1000	815	8.0	70.1	CF-57-9-35
	Inconel	70NaF-30BeF ₂	1000	815	6.0	52.6	CF-57-9-35
	Inconel	24LiF-53NaF-23BeF ₂	1000	815	5.0	43.8	CF-57-9-35
	Inconel	36LiF-49NaF-15BeF ₂	1000	815	3.0	26.3	CF-57-9-35
	Inconel	74LiF-26ThF ₄	1000	815	6.0	63.9	CF-57-9-35
517	Inconel	NaF-ZrF ₄	822	815	5.5	83.8	2338
337	Inconel	NaF-ZrF ₄	575	815	8.0	140.2	2338
214	Inconel	FLiNaK + NaK	500	815	3.0	52.6	1294
230	Inconel	36NaF-18KF-46ZrF ₄	500	815	10.0	175.2	1375
348	Inconel	50NaF-50ZrF ₄	500	815	5.5	96.4	1692
	Inconel	FLiNaK	500	815	7.0	122.6	1816
934	Inconel	60NaF-40ZrF ₄	500	815	5.0	87.6	2157
935	Inconel	60NaF-40ZrF ₄	500	815	5.0	87.6	2157
	Inconel	NaF-BeF ₂	500	815	10.0	175.2	[76]
	Inconel	LiF-NaF-BeF ₂	500	815	5.0	87.6	[76]

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246	Inconel	52NaF-48ZrF ₄	500	815	8.0	140.2	2337
262	Inconel	57NaF-43BeF ₂	500	815	9.0	157.7	2337
277	Inconel	50NaF-50ZrF ₄	500	815	5.0	87.6	2337
276	Inconel	NaF-ZrF ₄	500	815	8.0	140.2	2338
277	Inconel	NaF-ZrF ₄	500	815	4.0	70.1	2338
336	Inconel	NaF-ZrF ₄	500	815	6.0	105.1	2338
341	Inconel	NaF-ZrF ₄	500	815	5.5	96.4	2338
342	Inconel	NaF-ZrF ₄	500	815	6.0	105.1	2338
516	Inconel	NaF-ZrF ₄	500	815	6.0	105.1	2338
338	Inconel	NaF-ZrF ₄	500	815	6.0	210.2	2338
411	Inconel	NaF-ZrF ₄	250	815	4.5	394.2	2338
410	Inconel	NaF-ZrF ₄	100	815	4.0	700.8	2338
400	Inconel	NaF-ZrF ₄	50	815	3.0	3.0	2338
1181	Inconel	71LiF-29ThF ₄	8760	732	6.5	6.5	2684
1239	Inconel	71LiF-16BeF ₂ -13ThF ₄	8760	732	7.5	5.0	2973
9377-6	Inconel	71LiF-16BeF ₂ -13ThF ₄	13155	704	13.0	13.0	3215
1188	Inconel	35LiF-27NaF-38BeF ₂	8760	677	9.0	9.0	2723
1210	Inconel	71LiF-29ThF ₄	8760	677	5.0	5.6	2799
1235	Inconel	71LiF-16BeF ₂ -13ThF ₄	7789	677	4.0	7.5	2973
1214	Inconel	FLiNaK	4673	677	13.0	113.9	2684
1169	Inconel	71LiF-29ThF ₄	1000	677	1.0	8.8	2474
1177	Inconel	71LiF-29ThF ₄	1000	677	1.5	13.1	2474
1173	Inconel	58NaF-35BeF ₂ -7ThF ₄	1000	677	4.0	35.0	2474
1176	Inconel	58LiF-35BeF ₂ -7ThF ₄	1000	677	1.0	8.8	2474
1234	Inconel	71LiF-16BeF ₂ -13ThF ₄	1000	677	1.0	1.0	2799
9344-2	Inconel	FLiNaK	8760	649	8.0	8.0	2890
9344-2	Inconel	FLiNaK	8735	649	8.0	70.1	3215
1172	Inconel	35LiF-27NaF-38BeF ₂	1000	607	2.0	17.5	2474
1175	Inconel	FLiNaK	1000	607	1.0	2.9	2474
LDRD	INOR-8	FLiNaK	3048	815	0.1	0.1	[77]
1209	INOR-8	71LiF-29ThF ₄	8760	732	0.0	0.0	2799
1216	INOR-8	58LiF-35BeF ₂ -7ThF ₄	8760	732	1.0	1.0	2973
1240	INOR-8	71LiF-16BeF ₂ -13ThF ₄	8760	732	0.0	0.0	2973
MSRP	INOR-8	71LiF-16BeF ₂ -13ThF ₄	20000	704	1.0	0.9	3215

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7							
MSRP							
8	INOR-8	58LiF-35BeF ₂ -7ThF ₄	9633	704	0.0	0.0	3215
15A	INOR-8	73LiF-2BeF ₂ -25ThF ₄	39476	677	0.1	0.1	TM-4286
1208	INOR-8	FLiNaK	8760	677	1.0	1.0	2799
1190	INOR-8	58NaF-35BeF ₂ -7ThF ₄	8760	677	1.0	1.0	2799
1233	INOR-8	71LiF-16BeF ₂ -13ThF ₄	8760	677	0.0	0.0	2973
1213	INOR-8	71LiF-29ThF ₄	3114	677	0.0	0.0	2626
15	INOR-8	73LiF-2BeF ₂ -25ThF ₄	2003	677	0.0	0.0	TM-4286
1165	INOR-8	FLiNaK	1340	677	0.0	0.0	2551
1164	INOR-8	58NaF-35BeF ₂ -7ThF ₄	1000	677	0.0	0.0	2551
1221	INOR-8	71LiF-29ThF ₄	1000	677	0.0	0.0	2626
1228	INOR-8	71LiF-16BeF ₂ -13ThF ₄	1000	677	0.0	0.0	2723
MSRE	INOR-8	67LiF-33BeF ₂	26000	649	0.0	0.0	TM-4174
9354-3	INOR-8	35LiF-27NaF-38BeF ₂	19942	649	0.0	0.0	3215
1194	INOR-8	FLiNaK	1000	607	0.0	0.0	2551
1195	INOR-8	35LiF-27NaF-28BeF ₂	1000	607	0.0	0.0	2551

Note: The corrosion of austenitic iron and nickel based alloys such as Type 316 stainless steel and Inconel have been studies by Oak Ridge National Laboratory in a variety of high-temperature fluoride salts, with the results given in Table 12. Results for the alloy INOR-8 are also reported. Corrosion rates for the iron-based stainless steel appear to be bounded between approximately 5 and 150 mils per year. Relatively little corrosion data appears to exist under conditions relevant to the LIFE engine and must be collected. Testing is underway at LLNL to collect the required data [Williams et al. 2006].

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Table 16 – Solubilities of Chromium in Various Fluoride Salts

Salt mixture	ZrF ₄ or BeF ₂ (mol %)	UF ₄ (mol %)	Cr at 600°C (ppm)	Cr at 800°C (ppm)
FLiNaK	0	2.5	1100	2700
LiF-ZrF ₄	48	4	2900	3900
NaF-ZrF ₄	50	4.1	2300	2550
NaF-ZrF ₄	47	4	1700	2100
NaF-ZrF ₄	41	3.7	975	1050
KF-ZrF ₄	48	3.9	1080	1160
22NaF-55LiF-23ZrF ₄	23	2.5	550	750
LiF-BeF ₂	48	1.5	1470	2260

Note: As previously discussed, chromium from the ODS ferritic steel is expected to dissolve into both the FLIBE in the primary cooling loop, as well as into the FLiNaK in the secondary cooling loop. The dissolution of alloy chromium will be limited to some extent by the saturation concentration of the chromium in the fluoride salt. These limiting solubilities are given in Table 13 [Williams et al. 2006], and have been used to estimate the maximum wall penetration expected in the steel LIFE engine. The tabulated solubility for beryllium in fluoride salts further illustrates the facile dissolution of the neutron multiplier in these salts. Uranium is of course also readily dissolved, which is beneficial in the formation of damage tolerant liquid fuels.

Table 17 – Worldwide LWR SNF Inventories & Required YME Repositories

	LWR Plants	LWR Generating Capacity	LWR Total Stored SNF	Yucca Mountain Equivalent Needed	Cost for Yucca Mountain Equivalents
	#	GWe	kT HM	#	U.S. \$B
Western Europe	146	126	72	1.14	103
Eastern Europe	67	46	34	0.54	49
North America	124	112	105	1.67	150
Asia & Africa	104	75	33	0.52	47
World	441	359	244	3.87	349

Note: Data on worldwide fleet of light water reactors and spent fuel generated by those reactors was taken from K. Fukuda, W. Danker, J. S. Lee, A. Bonne, M. J. Crijns, IAEA Overview of Global Spent Fuel Storage, IAEA-CN-102/60, Department of Energy, International Atomic Energy Agency, Vienna, Austria, 2007, Table I. The assumed cost for a Yucca Mountain Equivalent (YME) repository was based upon recent statements by the United States Department of Energy: "DOE Testifies, Yucca's Costs Doubled, New Estimates of More Than \$90 Billion", Nuclear Waste News, Capitol Press LLC, Vol. 28, No. 15, July 21, 2008, p. 1. The Cost for the Yucca Mountain Repository has increased from the original life-cycle cost of \$58 billion to more than \$90 billion. The increased cost is due to a variety of factors, including the cost of temporarily storing the current of spent nuclear fuel, as well as escalating costs for construction and procurement of materials for the engineered barrier system. Given the escalating cost for geological disposal and such strong public and governmental opposition, any nuclear power-generation that can increase the quantity of electrical energy serviced by facility such as Yucca Mountain, thereby decreasing the need for a larger number of future repositories, should be welcome.

Table 18 – Equivalent Worldwide LIFE SNF Inventories & Required YME Repositories

	LIFE Engines	LIFE Generating Capacity	LIFE Total SNF	Yucca Mountain Equivalents Needed	Cost for Yucca Mountain Equivalents
	#	GWe	kT HM	#	U.S. \$B
Western Europe	60	126	4.3	0.07	6
Eastern Europe	22	46	2.1	0.03	3
North America	54	112	6.3	0.10	9
Asia & Africa	36	75	2.0	0.03	3
World	172	359	14.6	0.23	21

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Table 19 – Comparison of LWR & Equivalent LIFE Fleets: Required YME Repositories

	LWR YME Needed	LWR YME Cost	LIFE YME Needed	LIFE YME Cost	Cost Differential LWR-LIFE
	#	U.S. \$B	#	U.S. \$B	U.S. \$B
Western Europe	1.14	103	0.07	6	96
Eastern Europe	0.54	49	0.03	3	46
North America	1.67	150	0.10	9	141
Asia & Africa	0.52	47	0.03	3	44
World	3.87	349	0.23	21	327

Table 20 – Comparison of Mass Flow & TRU Generation for Various Fuel Cycles

	Reactor Scheme	Mass Flow of Heavy Metal		ompositio	on After I	rradiation	(Wt.% o	f IHM)
		kg/TWhe	U	Np	Pu	Am	Cm	Total TRU
1	LIFE 1-GWe DU fuel (99% FIMA)	137	0.5	0.004	0.07	0.02	0.44	0.53
	())/() I IIVIA)	137	0.5	0.004	0.07	0.02	0.44	0.55
2	LIFE 2-GWe DU fuel							
	(99% FIMA)	123	0.5	0.004	0.07	0.02	0.44	0.53
3	Once-through PWR (60							
	GWt-day/MTHM)	2050	98.5	0.10	1.35	0.08	0.01	1.54
4	PWR w/ Pu burning in							
	MOX (1-pass							
	reprocessing)	225	91.5	0.02	7.24	0.68	0.16	8.10
5	PWR w/ Pu+Np							
	burning in MOX (1-							
	pass reprocessing)	215	90.4	0.42	8.35	0.71	0.14	9.62
6	PWR w/ Pu burning in							
	EU-MOX (multi-pass							
	reprocessing)	575	89.5	0.05	9.32	0.89	0.22	10.48
7	PWR w/ Pu+Am							
	burning in EU-MOX							
	(multi-pass							
	reprocessing)	238	91.3	0.06	7.74	0.71	0.18	8.69
8	PWR w/ Pu+Am							
	burning in EU-MOX							
	(multi-pass							
	reprocessing)	711	92.7	0.09	6.05	0.84	0.34	7.32
9	DUPIC cycle (1-pass							
	reprocessing + burning	1997	99	0.06	0.87	0.04	0.01	0.98
	of spent PWR fuel in				•			

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	CANDU)							
10	European FR (fully closed cycle)	890	77.6	0.12	21.12	0.88	0.2	22.32
11	PWR + Am, Pu, (Cm) burning in FR (multi- pass reprocessing)	390	78.6	0.07	20.71	0.55	0.05	21.38
12	Double Strata System: PWR+FR+Accelerator transmutation	106	57.1	0.06	39.81	2.56	0.51	42.94
13	PWR+IFR (fully closed cycle)	289	69.8	0.65	26.6	2	0.98	30.23
14	Gen IV gas-cooled IFR (fully closed cycle)	849	79.3	0.15	19.48	0.87	0.24	20.74
15	Accelerator-driven transmutation of minor actinides	46	5.4	6.09	47.58	23.15	17.72	94.54
16	Accelerator-driven transmutation of TRU	117	1.9	3.29	73.48	12.37	8.96	98.10

Note: Taken from Advanced Nuclear Fuel Cycles and Radioactive Waste Management, NEA No. 5990, Nuclear Energy Agency, Organization for Economic Co-Operation and Development, 2006, p. 30, Table 2.2.

Table 21 – Assumed System Properties for Interim Storage

Container Inner Cylinder Volume (m ³)	V-inside	10.29	10.29
Heat Generation Per Volume (W/m ³)	A	5,092	5,092
Container Cylinder Area (m ²)	S_{od}	50.84	50.84
Energy Flux (W/m ²) - Based on Heat Generation	q_{od}	1030	1030.32
Energy Flux (W/m ²) - Air Heat Transfer Coefficient	q_{od}	1030	1030.32
Area Fraction Available for Heat Conduction	f	0.10	0.05
Fuel Thermal Conductivity (W/m-K)	k_{fuel}	87	87
Effective Fuel Thermal Conductivity (W/m-K)	k_{fuel}	8.7	4.35
Container Thermal Conductivitry (W/m-K)	k _{metal}	60	60
Radius - Inner Cylinder (m)	b	0.82	0.82
Radius - Outer Cylinder (m)	a	1.36	1.36005
Air Heat Transfer Coefficient (W/m2-K)	h_{air}	11.35	11.35
Air Temperature (K)	Tair	333	333
Air-Container Interface Temperature (K)	$T_a = T_{od}$	424	424
Container-Fuel Interface Temperature (K)	$T_b = T_{id}$	436	436
Maximum Fuel Temperature (K)	T _{max}	534	632
Container Cross Section (m ²)		5.81	5.81
Total Channel Cross Section (m ²)		17.41	17.41
Square Edge Dimension (m)		4.17	4.17

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Table 22 – Interim Storage Assuming Area Fraction Available for Heat Conduction = 0.10

Years	5	10	15	20	30	50	100	150	200	300
% 5-Year Power	100.00	72.33	52.74	40.67	31.61	24.67	7.55	2.57	1.04	0.40
Total Power (kW)	548	397	289	223	173	135	41	14	6	2
Container Power at 95% FIMA (kW)	52	38	28	21	17	13	4	1	1	0
Air Temp. (°C)	60	60	60	60	60	60	60	60	60	60
Air-Container Interface Temp. (°C)	151	151	151	151	151	151	151	151	151	151
Container-Fuel Interface Temp. (°C)	163	159	157	156	155	154	152	151	151	151
Fuel Centerline Temp. (°C)	261	231	209	196	186	178	159	154	152	151
Cooling Air (m ³ /min)	696	503	367	283	220	172	53	18	7	3
Air Velocity (m/sec)	1.0000	0.7233	0.5274	0.4067	0.3161	0.2467	0.0755	0.0257	0.0104	0.0040

Note: After removal from the LIFE engine, the fuels will be thermally and radioactively hot, and will require cooling to survive in interim storage and the repository. The characteristics of a hypothetical interim storage facility for LIFE spent nuclear fuel (SNF) are summarized in Tables 15 and 16. The temperature levels that will be experienced by the fuel materials during interim storage have been calculated, with the results summarized in Table 17 and 18 (two different heat transfer scenarios). With proper ventilation, the centerline of the stored spent fuel from the LIFE engine should not exceed 250 to 300°C. This temperature is well below the melting points of fuel materials, as well as the melting points of eutectics that are expected to form between silicon carbide encapsulation materials and fission products such as palladium. The Repository Design Team for LIFE continues to explore and optimize the interim storage and repository design for this hybrid fuel [Farmer et al. 2008].

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Table 23 – Interim Storage Assuming Area Fraction Available for Heat Conduction = 0.05

Years	5	10	15	20	30	50	100	150	200	300
% 5-Year Power	100.00	72.33	52.74	40.67	31.61	24.67	7.55	2.57	1.04	0.40
Total Power (kW)	548	397	289	223	173	135	41	14	6	2
Container Power (kW) - 95% FIMA	52	38	28	21	17	13	4	1	1	0
Air Temp. (°C)	60	60	60	60	60	60	60	60	60	60
Air-Container Interface Temp. (°C)	151	151	151	151	151	151	151	151	151	151
Container-Fuel Interface Temp. (°C)	163	159	157	156	155	154	152	151	151	151
Fuel Centerline Temp. (°C)	359	302	261	236	217	202	167	156	153	152
Cooling Air (m³/min)	696	503	367	283	220	172	53	18	7	3
Air Velocity (m/sec)	1.0000	0.7233	0.5274	0.4067	0.3161	0.2467	0.0755	0.0257	0.0104	0.0040

Table 24 – Key radionuclides in TSPA for Yucca Mountain

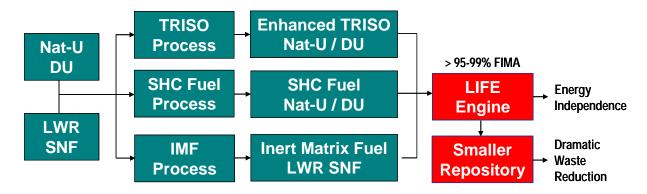
	< 10,000	10,000 to 100,000	100,000 to
	Years	Years	1,000,000
			Years
Most important radionuclides overall	⁹⁹ Te, ¹⁴ C, ¹²⁹ I, ²³⁹ Pu	²³⁹ Pu, ¹²⁹ I, ²²⁶ Ra	²²⁶ Ra, ²⁴² Pu, ²³⁷ Np
Other nuclides important for specific scenarios	²⁴⁰ Pu, ²⁴¹ Am	⁷⁹ Se, ⁹⁹ Tc, ¹³⁵ Cs, ²³⁷ Np, ²⁴⁰ Pu, ²⁴² Pu	⁷⁹ Se, ⁹⁹ Tc, ¹³⁵ Cs, ²³⁷ Np, ²⁴⁰ Pu, ²⁴² Pu

Note: Nuclides identified in the Yucca Mountain TSPA-LA as dominating the doses for different time periods from a Yucca Mountain repository [Shaw et al. 2008].

Figures Fissile Molten Salt Fuel Direct > 80-99% FIMA Material WG-Pu / HEU **Digestion** Incineration **Prototype** WG-Pu **Engine** HEU **TRISO Enhanced TRISO** Initial LIFE **Process** WG-Pu / HEU Demo

There are three primary missions for the LIFE engine, which include the burning of excess weapons-grade plutonium (WG-Pu), natural and depleted uranium (DU and HEU), and light-water reactor (LWR) spent nuclear fuel (SNF). As shown here, WG-Pu and HEU can be digested directly into a molten salt, thereby forming a high burn-up liquid fuel for the direct incineration of old weapons materials. TRISO = TRI-structural ISO-tropic; SHC = Solid Hollow Core; IMF = Inert Matrix Fuel; FIMA = Fission of Intial Metal Atoms (burn-up) [Farmer et al. 2008].

Figure 1 – Mission for burning weapons grade uranium in LIFE engine.



Natural and depleted uranium (Nat-U and DU) can be incorporated into a variety of solid fuels, including TRI-ISO-tropic fuel (TRISO); baseball or solid hollow core (SHC), and inert matrix fuel (IMF). The LIFE engine enable the direct burning of such fuels with no isotopic enrichment or chemical separation, with the potential to attain very high burn-up, which is quantified in terms of the fraction initial uranium metal atoms (FIMA) burned [Farmer et al. 2008].

Figure 2 – Mission for natural and depleted uranium, and spent nuclear fuel from light-water reactors in LIFE engine.

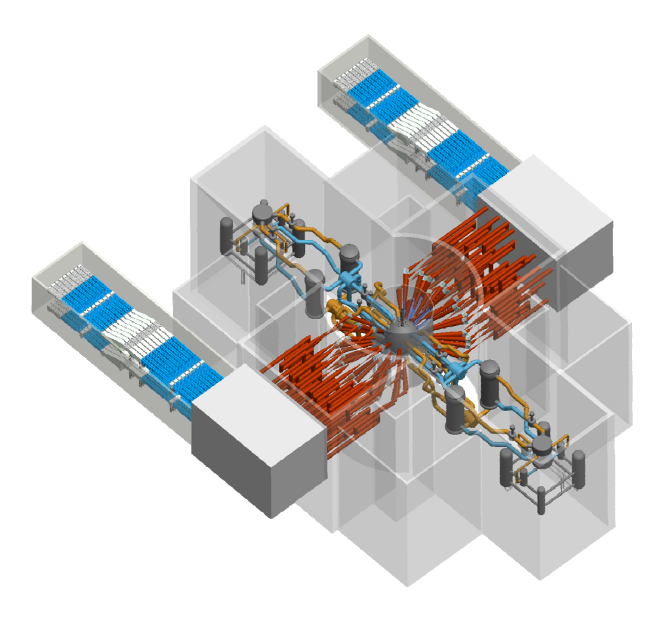


Figure 3 – LIFE plant with NIF-like central hot spot ignition.

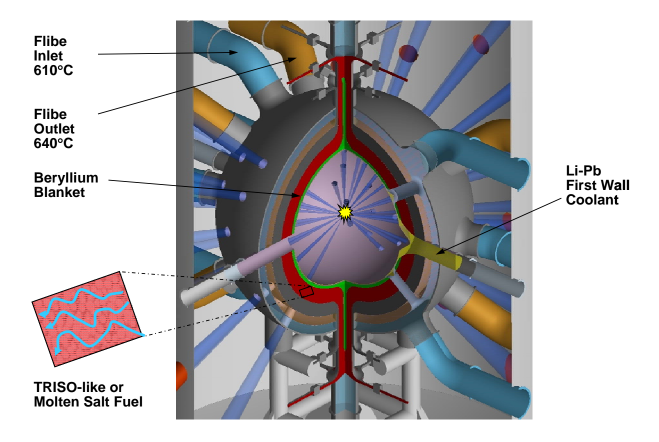


Figure 4 – LIFE engine with NIF-like central hot spot (CHS) inertial confinement fusion (ICF) target features a dedicated first wall coolant with a pebble-based multiplier and fuel.

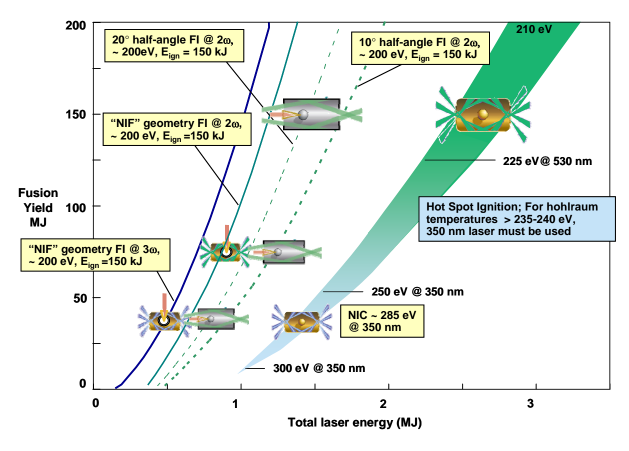


Figure 5 – There is a wide region of potential ICF target-laser options for LIFE.

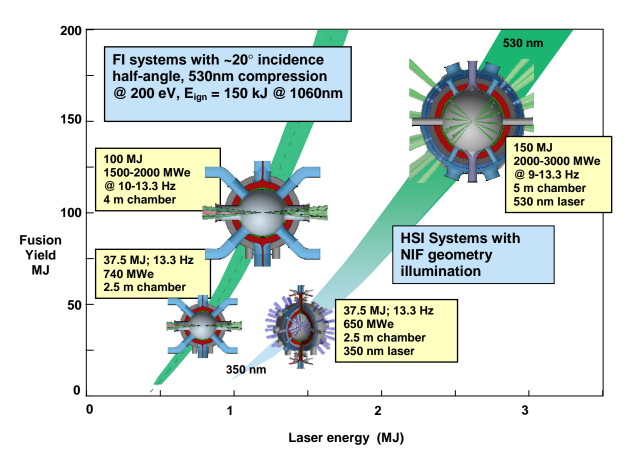


Figure 6 - Different targets would result in different chamber sizes and electric output.

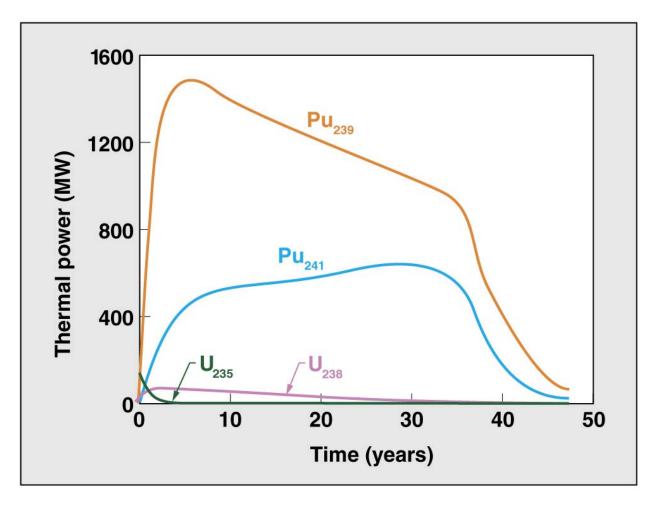


Figure 7 – When burning DU, several isotopes will undergo fission in the LIFE engine.

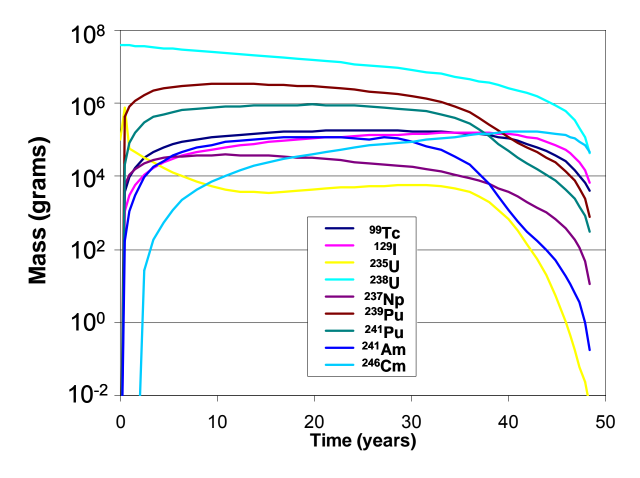


Figure 8 – LIFE fuel burn-up can be adjusted as desired.

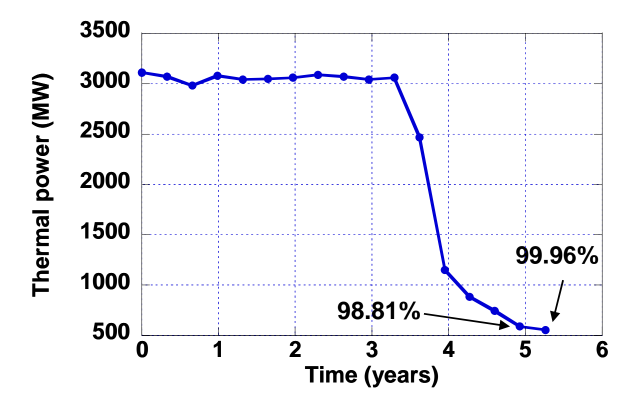


Figure 9 – LIFE can burn excess plutonium in a high-gain blanket. This conceptual system is assumed to be fueled with 7 tons of weapons-grade plutonium (WG-Pu). It is further assumed that: TRISO kernels are blended 80% ZrC + 20% Pu; TRISOs are loaded with 400 wppm boron as burnable poison; fusion power is 375 MW (37.5 MJ @ 10 Hz); the flat-top thermal power is 3000 MW; and the blanket gain is eight (8).

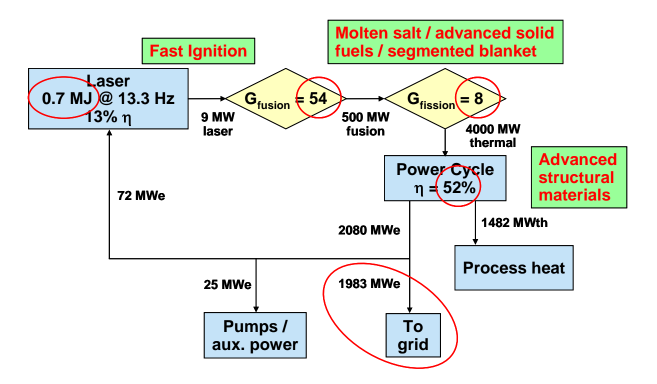


Figure 10 – An early LIFE demonstration unit, burning WG-Pu, will enable the development of an advanced power system capable of helping provide much of the Nation's future energy needs, with substantial reductions in the generation of high-level radioactive waste. Specific system improvements that are anticipated include the advent of fast-ignition ICF, advanced solid and molten salt fission blankets, and advanced structural materials capable of operating at higher temperatures.

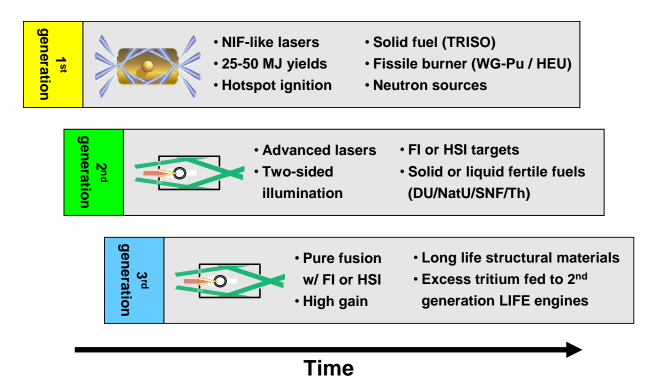
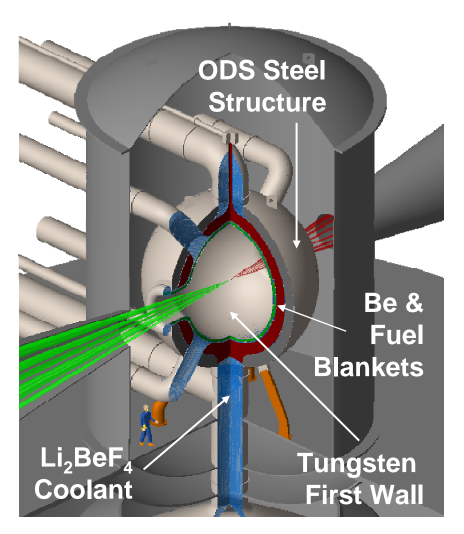
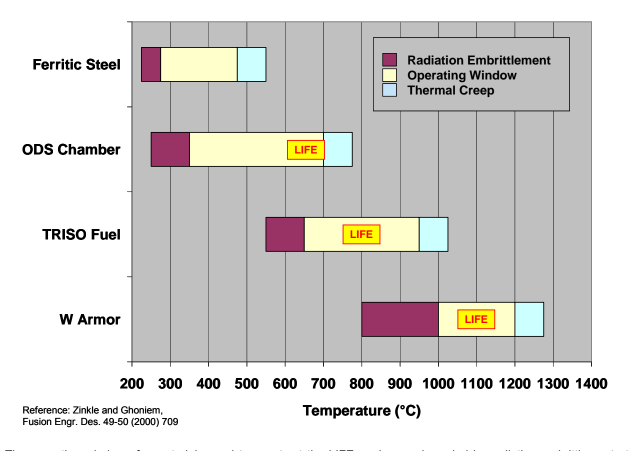


Figure 11 – The WG-Pu mission has the potential of enabling the hybrid LIFE systems, and will also enable pure fusion, since LIFE is a practical and realizable precursor for pure fusion.



LIFE engine's structural challenges include: need for high-temperature strength; resistance to high-temperature creep; immunity to radiation damage, including swelling and helium embrittlement; resistance to corrosion and environmental cracking in high-temperature molten fluoride salts; and the ability to be fabricated into necessary shapes and configurations with practical welding processes. These challenges are being met by ODS ferritic steels, refractory metal sheets and coatings of tungsten, advanced TRISO fuels, and other advanced reactor materials.

Figure 12 – LIFE engine with low incidence angle (LIA) fast ignition (FI) target, showing key materials: tungsten first wall, oxide dispersion strengthened steel structure, beryllium neutron multiplication blanket, fission blanket, and molten fluoride salt coolant. In contrast to the engine shown in Figure 4, compression beams enter from one side of the engine, and ignition beams enter from the opposite side.



The operating windows for materials used to construct the LIFE engine are bounded by radiation embrittlement at lower temperatures, and by thermal creep at higher temperatures. The operating conditions for LIFE have been selected to enable the survivability of known materials of construction.

Figure 13 – Temperature limits for LIFE engine materials of construction.

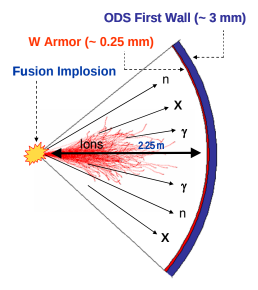


Figure 14 – Materials will be subjected to a severe radiation and high-temperature environment. Energy from ICF target carried by: neutrons (14 MeV); Ions (< 100 keV); X-rays (< 100 keV); gamma ray contribution small.

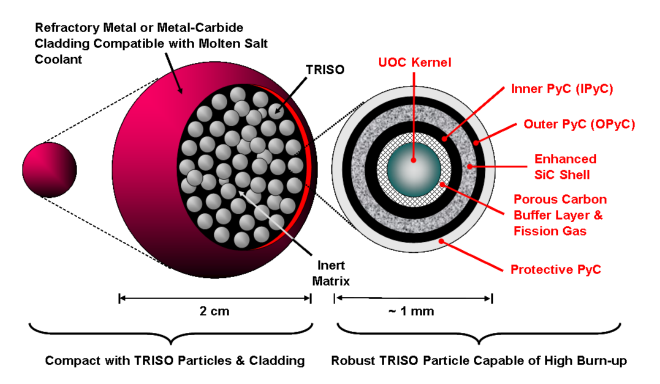
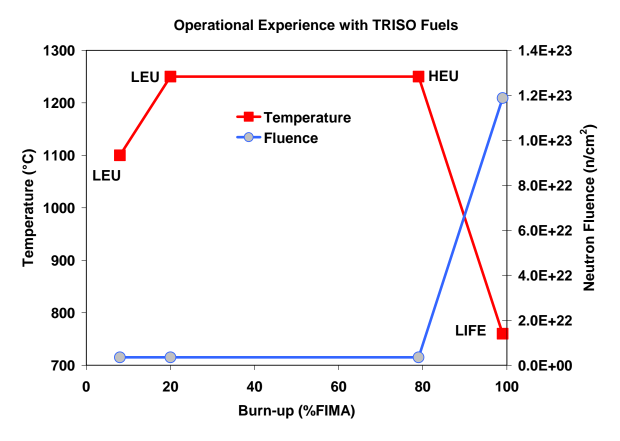
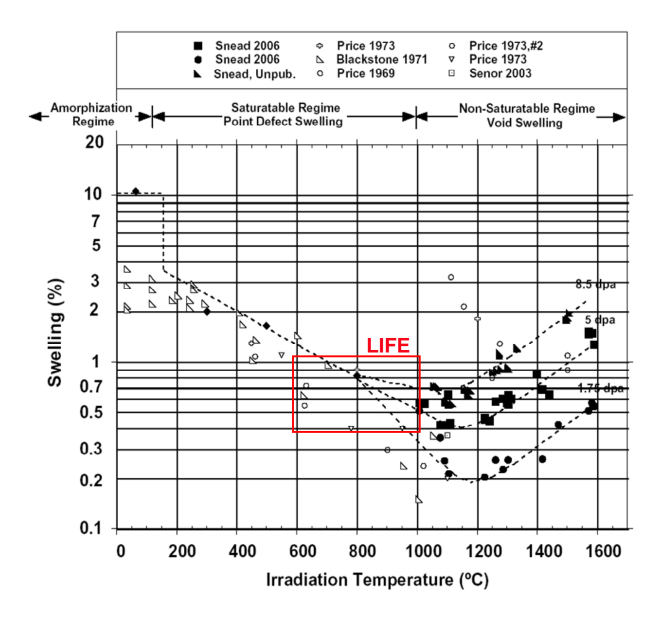


Figure 15 – An enhanced TRISO fuel, with a more robust SiC capsule to enable fission-gas containment, is being considered as one possible high burn-up fuel option by LIFE engine designers.



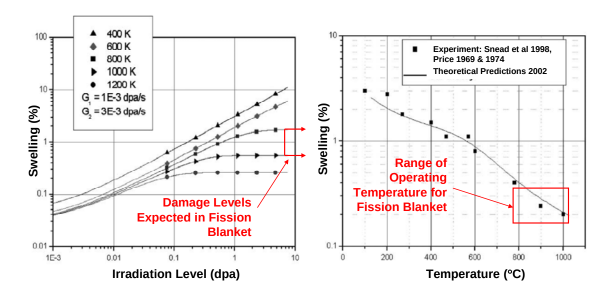
The experience base for TRISO fuels is limited, with 8 to 20% FIMA demonstrated with low enriched uranium (LEU), and 79% FIMA demonstrated with high enriched uranium (HEU). These LEU and HEU fuels experienced only $\sim 3.5 \times 10^{21}$ n/cm², but at a relatively high temperature of 1100 to 1200°C. In contrast, LIFE fuels will experience a much higher neutron dose of approximately 1.2×10^{23} n/cm², but at a much lower temperature of about 700°C.

Figure 16 – Experience with TRISO fuels compared to anticipated conditions in LIFE engine.



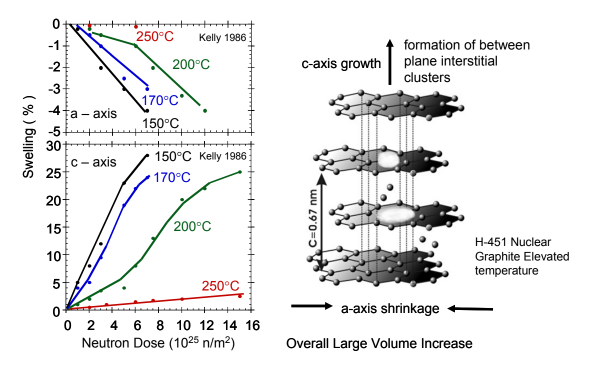
LIFE TRISO fuel will operate below 1000°C, avoiding the non-saturable void-swelling regime. Based upon published data for damage levels up to 8.5 displacements per atom (dpa), the swelling of SiC capsule expected to be 0.4 to 0.9 % [Snead et al. 2007]. Data at much higher irradiation levels is desperately needed.

Figure 17 – The fission blanket is expected to operate in saturable point-defect swelling regime of silicon carbide (SiC).



Based upon a detailed kinetic model that accounts for the growth of dislocation loops and ion-beam irradiation studies at Kyoto University's DuET facility, irradiation-induced swelling of SiC is expected to reach an asymptotic value of less than approximately one percent (~1%) as the dose increases [Ryazanov et al. 2002].

Figure 18 – Model predictions and ion-beam irradiation of SiC show no significant swelling of above approximately one displacement per atom (~ 1 dpa).



The swelling of the lattice normal to the hexagonal planes (along c-axis) is substantially greater than that parallel to the planes (along a-axis). A material that swells isotropically, and to a lesser extent, is preferable [Zinkle 2008].

Figure 19 – The anisotropic swelling of the inner and outer pyrolytic graphite (IPyC and OPyC) layers in TRISO particles, as well as the graphite binder in the pebbles, is considered to be a significant barrier to the development of high burn-up TRISO fuels.

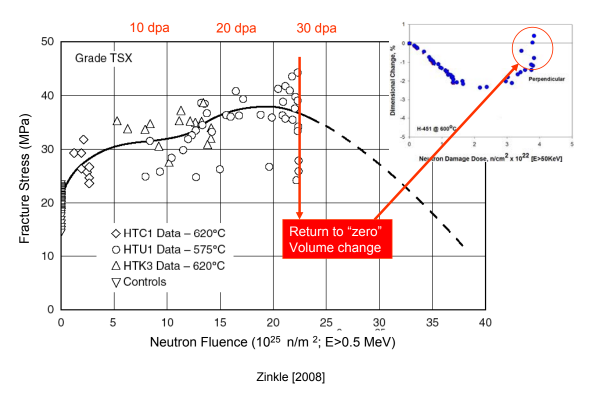
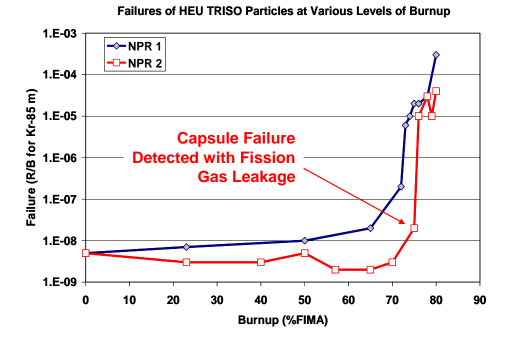
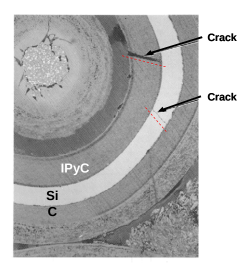


Figure 20 – The fracture strength of irradiated nuclear graphite is expected to be 25 to 45 MPa at $20 \text{ to } 25 \times 10^{25} \text{ n/m}^2$, but may drop at higher doses.



The observed probability of failure appeared to be approximately 1 in 77,500 at this level of burn-up. Achieving such high burn-up with TRISO fabricated with natural or depleted uranium is a much greater challenge [Petti et al. 2004].

Figure 21 – TRISO fabricated from HEU has achieved relatively high burn-up (79% FIMA) with our failure of the SiC capsule and leakage of fission gas such as Kr-85.



The observed leakage of fission gas from these particles appears to be due to the pressurization-driven stress and associated fracture of the spherical silicon carbide pressure vessel [Petti et al. 2004].

Figure 22 – This figure shows fractures in the IPyC and SiC layers of a TRISO particle after reaching 79% FIMA.

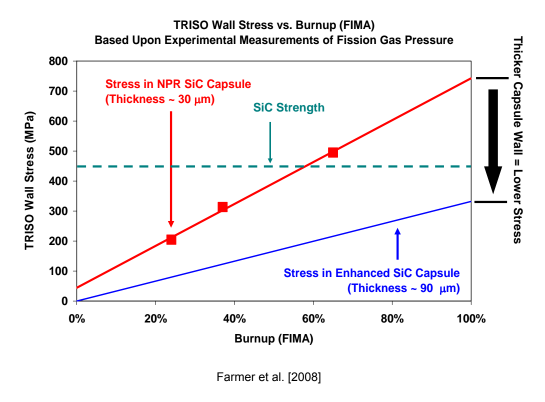
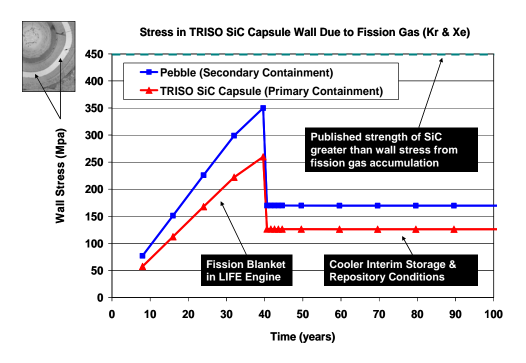
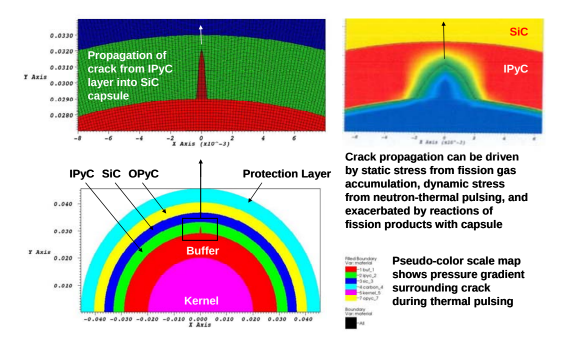


Figure 23 – The failure of these TRISO particles appears to coincide with the point where the SiC wall stress exceeds the ultimate tensile stress (UTS) of the material. One method of mitigating the failure is to thicken the SiC wall.



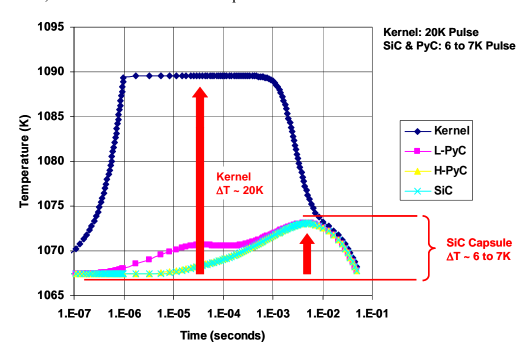
TRISO particles for LIFE engine are designed with enough silicon carbide wall thickness for fission gas containment at 99.9% FIMA [Farmer and Zhao 2008].

Figure 24 – SiC wall stress verses fuel burn-up for enhanced TRISO fuel for LIFE engine.



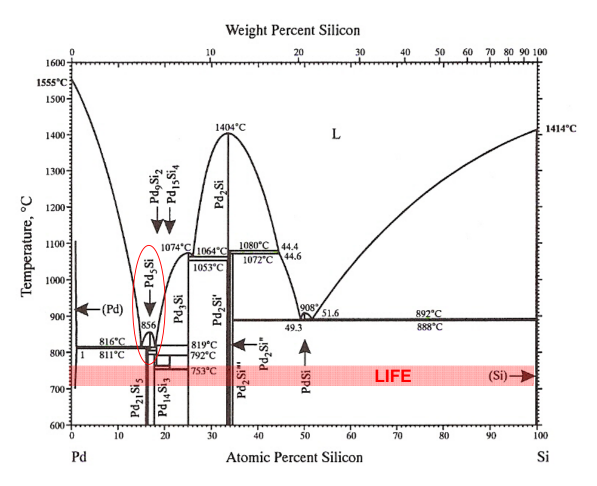
Model and calculations were done by Paul Demange et al. at LLNL [2008].

Figure 25 – Crack initiation at defects in silicon carbide layer are being accounted for in finite element model, and the calculations based upon that model.



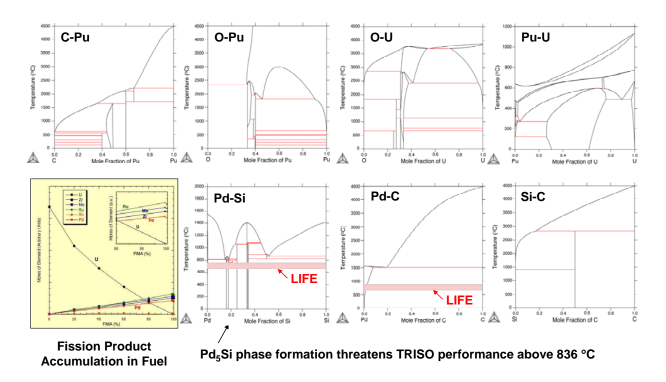
The temperature pulse in the fuel particle was modeled by Ryan Abbot at LLNL [2008].

Figure 26 – TRISO will experience dynamic stress due to thermal pulsing at 10 ot 20 Hz, which may cause thermal fatigue.



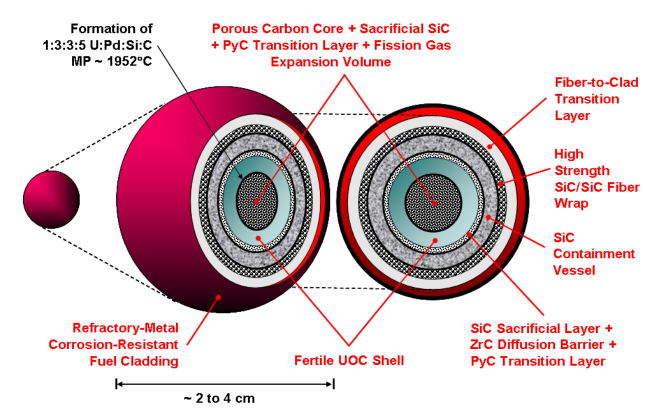
Survival of the SiC capsule is complicated by reactions involving fission products such as Pd. Deleterious low-melting eutectics form between fission products and encapsulation materials. A frequently cited example: is the Pd-Si reaction. Pd_5Si phase formation threatens TRISO performance above 836°C. Thermal migration of kernal in temperature gradient can bring such fission products in close proximity of the SiC. Mitigation includes the use of sacrificial SiC in (or around) the kernel and a ZrC diffusion barrier. The TRISO operating temperature in LIFE is below the melting point of deleterious phases such as Pd_5Si [Okamoto 2000].

Figure 27 – Phase diagram showing formation of low-melting Pd₅Si as a result of the reaction of a Pd (fission product) with SiC capsule in TRISO fuel particle.



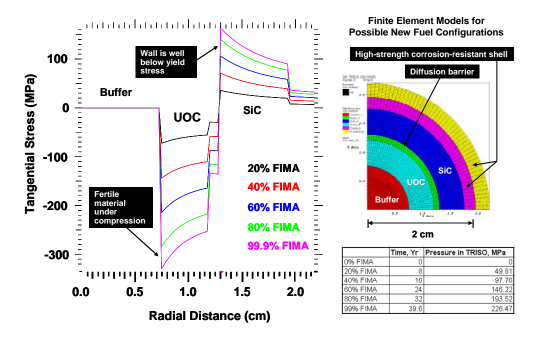
Kaufman and Turchi are beginning to use the THERMOCALC code to model phase transformations in fission fuels at very high burn-up [2008].

Figure 28 – Phase transformations in fission fuels at high burn-up are being modeled. Relevant binaries include: C-Pu, O-Pu, O-U, Pu-U, Pd-Si, Pd-C, Si-C & others.



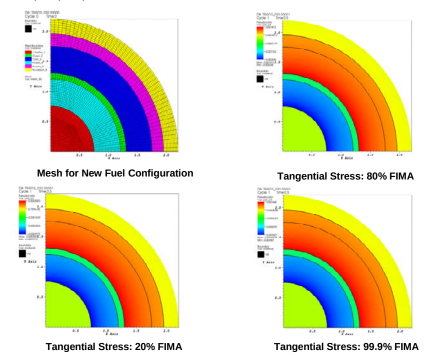
A wide-variety of alternative materials and fuel designs are possible and are being evaluated. Higher mass fraction of fertile material can be achieved with new solid hollow core (SHC) fuel [Farmer, Latkowski & Abbott 2008]. The specific use of sacrificial SiC in fuels is attributed to R. J. Lauf et al. [1984].

Figure 29 – Solid-hollow core (SHC) fuel design with high uranium mass fraction and enhanced strength encapsulation materials.



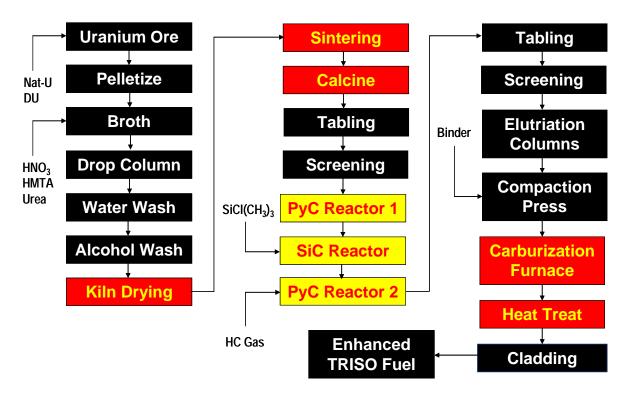
Model development and calculations were done by Paul Demange et al. at LLNL [2008].

Figure 30 – Predicted stress in the wall of solid hollow-core (SHC) fuel at fission gas pressures corresponding to 20, 40, 60, 80 and 99.9% FIMA.



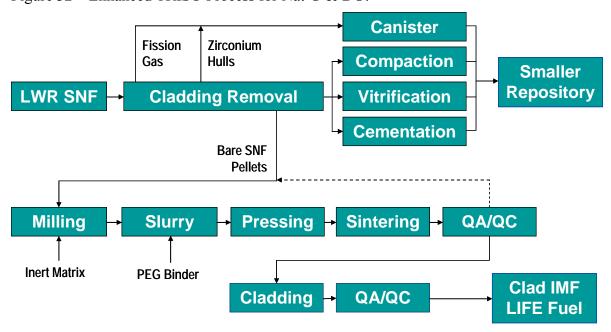
Model development and calculations were done by Paul Demange et al. at LLNL [2008].

Figure 31 – Numerical analysis indicates a substantial strength margin in SHC pressure boundary. In all cases, the stress is well below the yield and ultimate tensile stresses of the wall material.



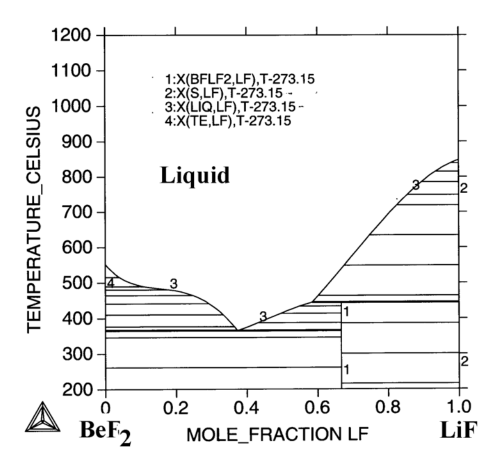
This conceptual process was developed by Farmer et al. at LLNL [2008].

Figure 32 – Enhanced TRISO Process for Nat-U & DU.



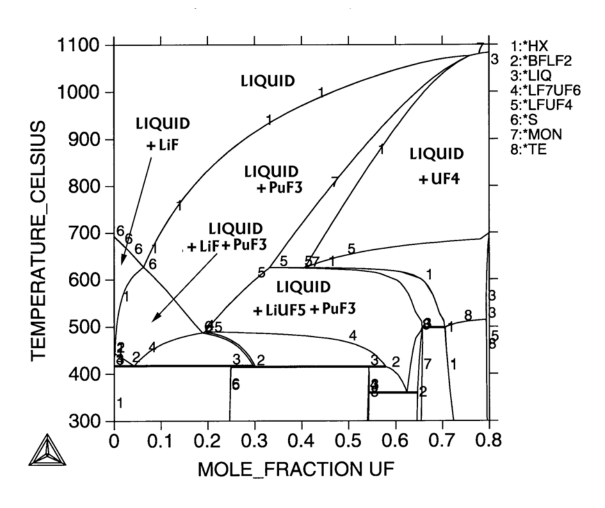
This conceptual process was developed by Farmer et al. at LLNL [2008].

Figure 33 – Other processes may be needed for the conversion of light water reactor spent nuclear fuel into LIFE fuel.



This molten salt coolant is less prone to fire than liquid metal coolants, and can be used to breed tritium from the lithium. The corrosion of structural components of the LIFE engine in contact with this coolant must be understood [Kaufman and Turchi 2008].

Figure 34 – A binary mixture of BeF₂ and LiF to form Li₂BeF₄ produces a eutectic composition with a low melting point of less than 400°C.



Molten salt fuel is not damaged by radiation and enables direct digestion of fertile/fissile materials. Phase transformations and actinide solubility in LIFE coolants and liquid fuels: BeF₂-LiF, BeF₂-ThF₄, BeF₂-UF₄, BeF₂-ZrF₄, LiF-PuF₃, LiF-ThF₄, LiF-ZrF₄, ThF₄-UF₄ and higher order systems. Regions of stable operation revealed for liquid fuels in LIFE engine [Kaufman and Turchi 2008].

Figure 35 – Phase diagram reveals regions of stable operation for liquid fuels in LIFE engine.

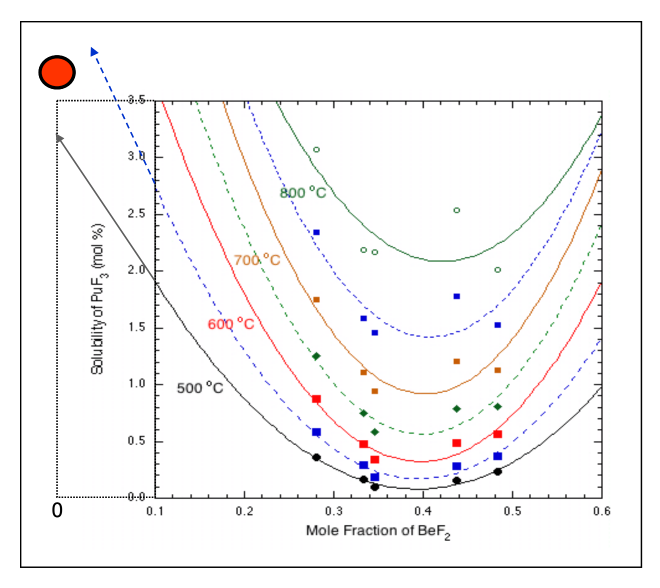


Figure 36 – Additional data will be required to determine whether or not a solubility of 2 mol% PuF_3 can be achieved at $550^{\circ}C$. While extrapolations have been used to suggest that PuF_3 may have solubility >3.5% at $550^{\circ}C$, data at lower concentrations of BeF_2 are needed.

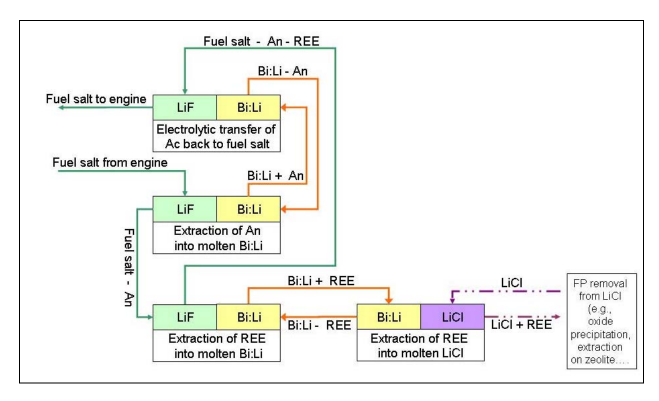
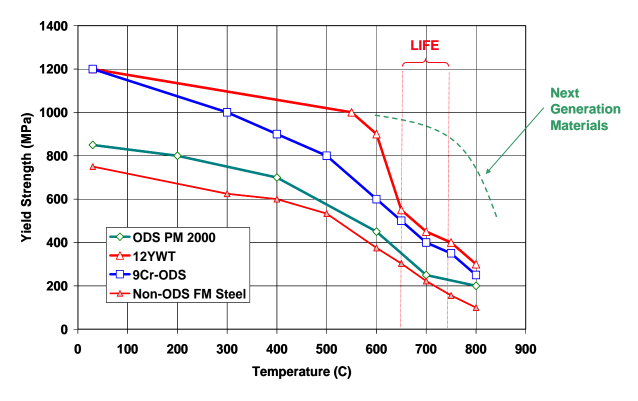
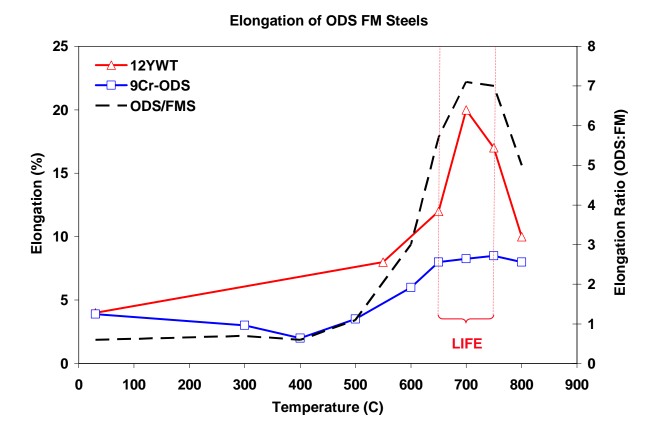


Figure 37 – Reductive extraction system for removal of rare earth fission products from molten salt fuel into metal. Continuous removal of rare earth fission products needed to prevent precipitation of plutonium tri-fluoride (PuF₃).



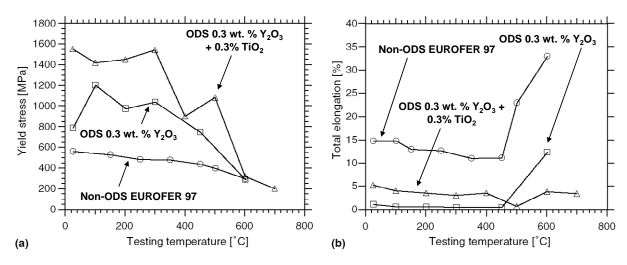
New ODS FM steels such as 9Cr-ODS & Kobe 12YWT have demonstrated exceptional tensile strength at high temperature; additional development is needed. ODS steels have exceptional tensile strength at high temperature; the next generation is expected to be better [Ukai and Fujiwara 2002; Klueh et al. 2007].

Figure 38 – New ODS FM steels such as 9Cr-ODS & Kobe 12YWT have demonstrated exceptional tensile strength at high temperature.



New ODS FM steels such as 9Cr-ODS & Kobe 12YWT have demonstrated exceptional creep resistance at high temperature; additional development is needed. ODS steels have exceptional tensile strength at high temperature; next generation is expected to be better [Ukai and Fujiwara 2002; Klueh et al. 2007].

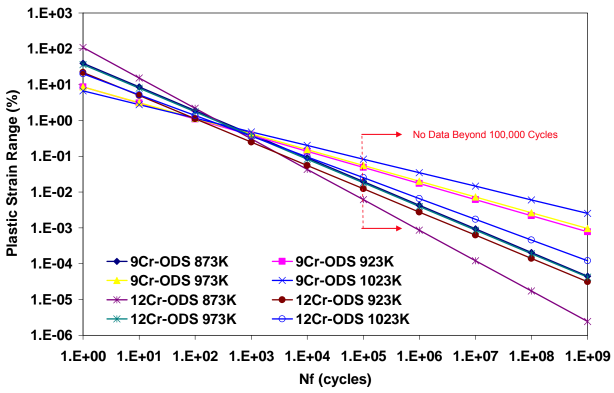
Figure 39 – New ODS FM steels such as 9Cr-ODS & Kobe 12YWT have demonstrated exceptional creep resistance at high temperature; additional development is needed.



Oxide dispersion strengthened (ODS) steels are the most promising class of materials with the potential to be used at elevated temerpature under severe neutron exposure. Leading technology development of these mateirals has been conducted at JNC with an emphasis on cladding for fast reactors. Tensile yield strength (a) and total elongation (b) as function of test temperature of (-O-) as-received EUROFER 97, (- \Box -) CRPP ODS EUROFER97 0.3 wt. % Y₂O₃ and (- \Box -) CRPP ODS EUROFER97 0.3 wt. % Y₂O₃ + 0.3 wt. % TiO₂ [Schaublin et al. 2006].

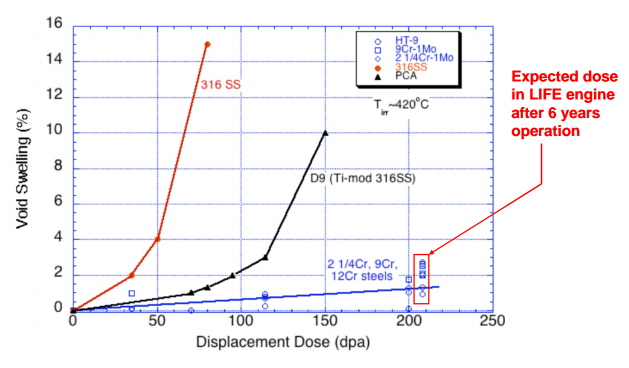
Figure 40 – Additional data for EUROFER 97 showing the exceptional high-temperature strength and creep resistance of ODS steels.

Plastic Strain Behavior of 9Cr-ODS & 12Cr-ODS



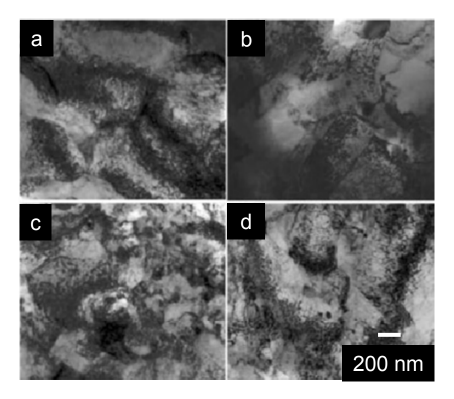
ODS ferritic-martensitic steel has longer fatigue life than conventional ferritic steels such as modified 9Cr-1Mo seel. Predictions based upon regression analysis of data for N_f < 100,000 cycles. Strain-controlled low-cycle fatigue tests were conducted for 9Cr-ODS and 12Cr-ODS at 873-1023K (600 to 800°C). The total strain ranges were controlled from 0.5 to 1.5% with a strain rate of 0.1% per second. Corresponding plastic strain ranged from 0.01 to 1%. The ODS has longer fatigue life than modified 9Cr-1Mo ferritic steel. The oxide dispersion strengthening prevented softening or hardening during cycling [Ukai and Ohtsuka 2007].

Figure 41 – Fatigue cycling of 9Cr-Fe and 12Cr-Fe ODS at temperatures from 600 to 800°C.



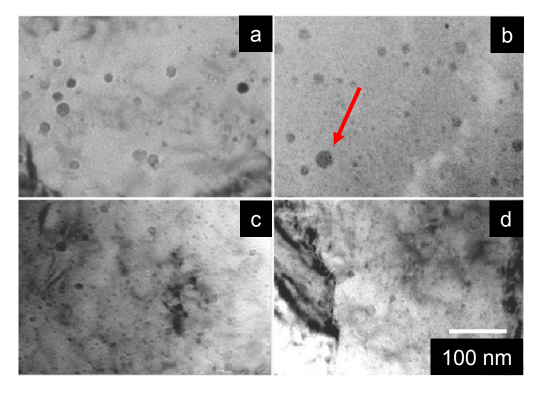
Gelles [1996]; Toloczko [2000]; Klueh and Harries [2001]; Zinkle [2008]

Figure 42 – Ferritric steels experience less irradiation-induced swelling than Type 316 austenitic stainless steel.



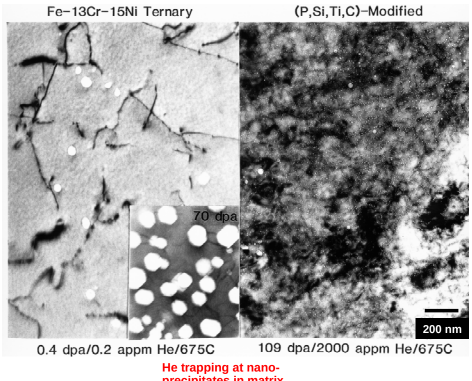
Overview of the microstructure in 9Cr-ODS irradiated with Ni ion at 500°C: (a) un-irradiated, (b) 5 dpa, (c) 50 dpa and (d) 150 dpa. No voids were found in all the conditions [Allen et al. 2005].

Figure 43 – Ion-beam irradiation of 9Cr-Fe ODS ferritic steel has shown no detectable void formation up to 150 displacements per atom.

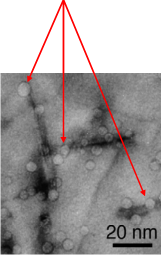


lon-beam irradiation of ODS steel has shown no detectable void formation up to 150 dpa. Oxide particles in 9Cr-ODS irradiated with Ni ion at 500°C: (a) un-irradiated, (b) 5 dpa, (c) 50 dpa, and (d) 150 dpa. The average size decreased from 11.7 nm for the un-irradiated case to 9.3 nm for the irradiated case of 150 dpa [Allen et al. 2005]

Figure 44 – Ion-beam irradiation of ODS steel has shown no detectable void formation up to 150 displacements per atom.



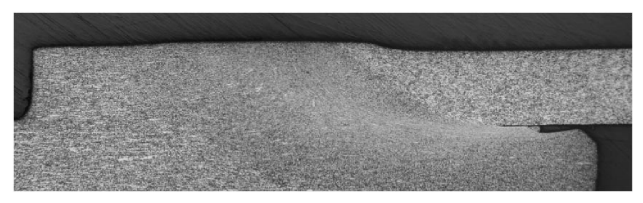
He trapping at nanoprecipitates in matrix



Dispersions of nano-precipitates mitigate irradiation-induced swelling attributable to helium. Helium trapping was observed at nano-precipitates in the matrix [Mansur and Lee 1991].

Figure 45 – Dispersions in matrix mitigate irradiation-induced swelling attributable to helium.

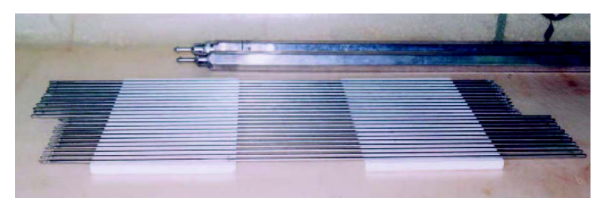
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Friction stir, inertial and pressurized resistance welding provide best results Y_2O_3 particles remain in weld. Longitudinal cross-section in the vicinity of welded 9Cr-ODS steel cladding and its end-plug joined with pressurized resistance welding [Ukai et al. 2005].

Figure 46 – ODS FM steels have been successfully joined using industrial welding processes.

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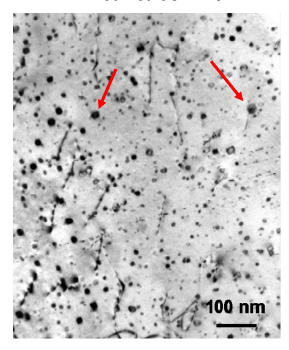




Fuel pins with ODS steel cladding and welded end-caps, joined with pressurized resistance welding, are being tested in BOR-60 reactor: 5% FIMA & 25 dpa by 2004; and 15% FIMA & 75 dpa by 2008. No failures of base metal or welds have been observed as 75-dpa level approached [Ukai et al. 2005].

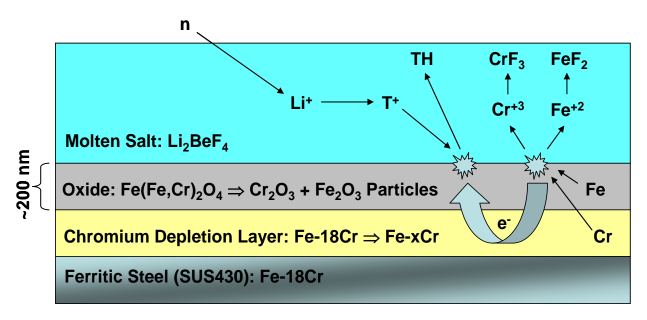
Figure 47 – Fuel pins with ODS steel cladding and welded end-caps.

TMT Modified 9Cr-1Mo



Other approaches to dispersion-strengthening also exist, such as thermomechanical treatment (TMT). This process uses hot rolling to process 9Cr-1Mo and 12 Cr steels. Precipitates are formed at dislocations introduced during such processing. Dispersions of precipitates produced with TMT are much finer than those observed in conventionally processed 9-12 Cr steels. Mechanical properties are substantially improved. [R.L. Klueh et al. 2005].

Figure 48 – The microstructure of 9-12% Cr steels can be improved dramatically with hot rolling, which is known in the literature as thermomechanical treatment (TMT).



Note: The corrosion mechansim for ODS ferritic steel is complicated, as illustrated in this figure. First, the transmutation of lithium in the Li_2BeF_4 salt leads to the formation of acidic tritium fluoride, which has the chemical behavior of hydrofluoric acid. Based upon discussion in the literature, it is believed that gaseous beryllium containing species in equilibrium with the molten salt electrolyte may include $LiBeF_3$, BeF_2 and BeH_2 . The reduction of T^+ at the ODS-salt interface may lead to the formation of reduced T_2 , with the simultaneous oxidation of Fe and Cr in the alloy to form Cr^{+3} and Fe^{+2} . Atomic tritium adsorbed on the surface of the ODS steel could be absorbed by the steel, which could lead to tritium (hydrogen) induced cracking. The preferential dissolution of chromium into the Li_2Be_4 salt results in the formation of a chromium depletion layer, which lies below an oxide layer, consisting of a $Fe(Fe,Cr)_2O_4$, with an layer of dispersed Fe_2O_3 particles [Farmer 2008].

Figure 49 – Mechanism for corrosion of ODS ferritic steel, showing formation of the chromium depletion layer, and dissolution of iron and chromium into the molten salt.

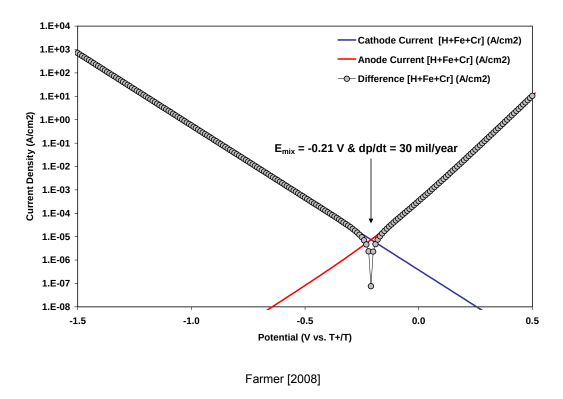


Figure 50 – Mixed potential theory applied to ODS ferritic steel without sacrificial beryllium anode in molten Li₂BeF₄ at 815°C.

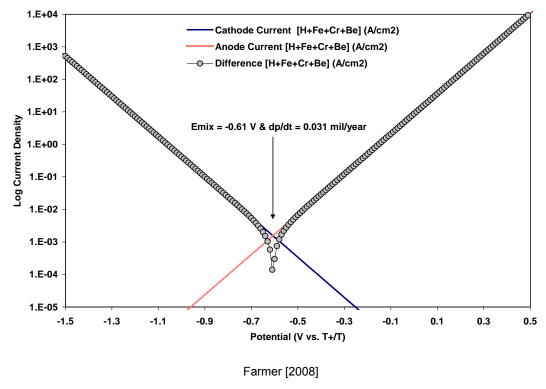


Figure 51 - Mixed potential theory applied to ODS ferritic steel with sacrificial beryllium anode in molten Li_2BeF_4 at 815°C .

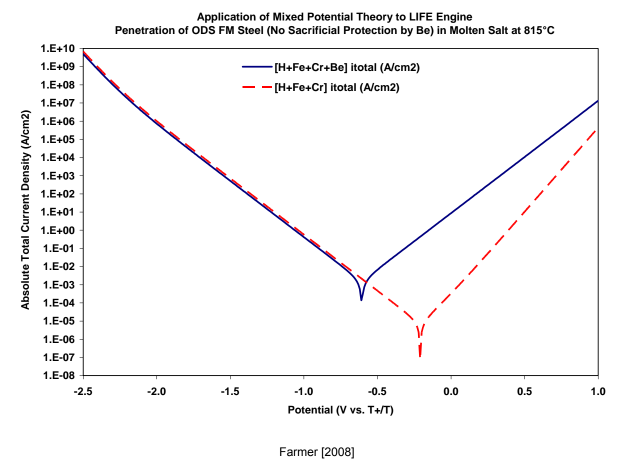


Figure 52 - A comparison of these two curves, taken from Figures 39 and 40 respectively, clearly show the cathodic shift of the open circuit corrosion potential due to galvanic coupling of beryllium and ODS steel at 815° C.

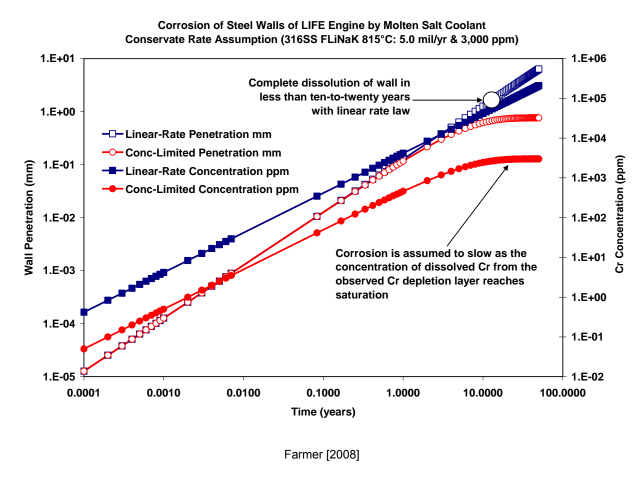


Figure 53 – Limited corrosive attack of ODS by molten salts, assuming a rate of attack of 5.0 mils per year, and saturation of the salt with chromium at a concentration of approximately 3,000 parts per million.

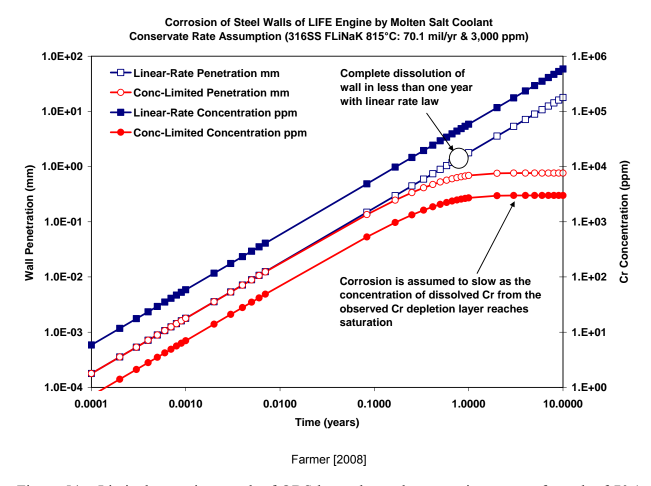
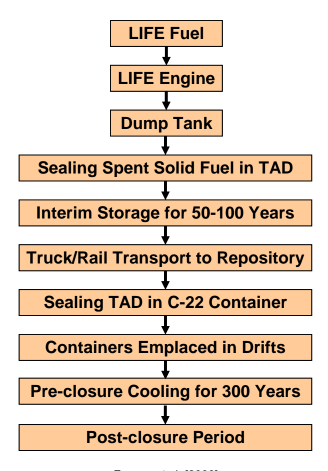


Figure 54 – Limited corrosive attack of ODS by molten salts, assuming a rate of attack of 70.1 mils per year, and saturation of the salt with chromium at a concentration of approximately 3000 parts per million.



Farmer et al. [2008]

Figure 55 – LIFE engine's fuel-cycle story, including interim storage and disposal of waste in geological repository.

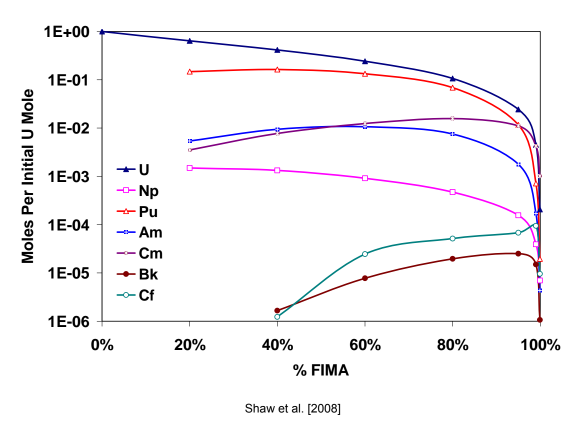


Figure 56 – TRU inventory in fission blanket of LIFE engine (spent fuel) at various levels of burn-up (FIMA).

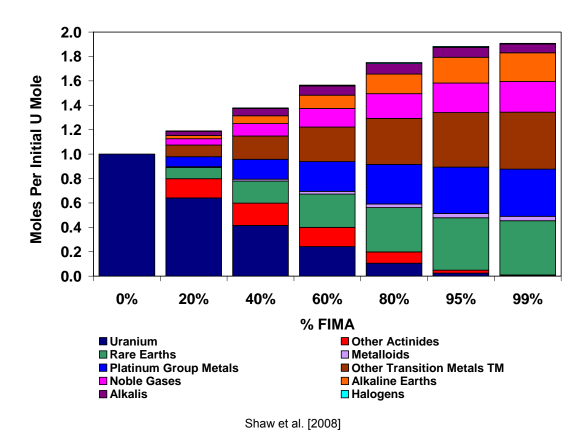
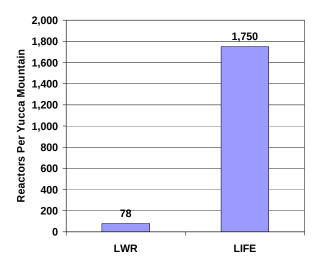
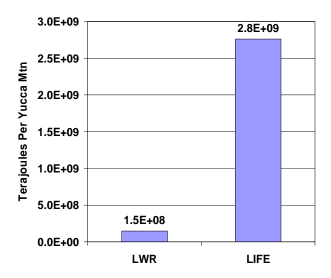


Figure 57 – Compositon of fission blanket of LIFE engine (spent fuel) at various levels of burn-up (FIMA).



LIFE could extend the useful service life of deep geologic repositories by factor of twenty (×20). LIFE would require less repository capacity than LWR. Twenty times more electricity could be produced with nuclear before filling up repository [Farmer et al. 2008].

Figure 58 – Assuming current statutory limits for the disposal of spent nuclear fuel, a single Yucca Mountain repository could service more LIFE engines that light-water reactors.



LIFE could extend the useful service life of deep geologic repositories by factor of twenty (\times 20). Each reactor is assumed to have approximately 50-year service life: LWR = 1.2 GWe per 900 metric tons of heavy metal waste; LIFE = 1.0 GWe per 40 metric tons of heavy metal (MTHM) waste [Farmer et al. 2008].

Figure 59 – A single Yucca Mountain repository could enable more electrical power generation with a fleet of LIFE engines than with a fleet of light-water reactors.

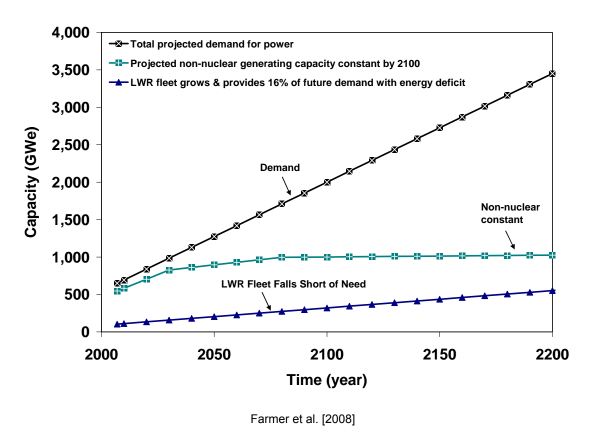


Figure 60 – Electrical generating capacity for 1st Scenario: LWR fleet with no LIFE and energy deficit.

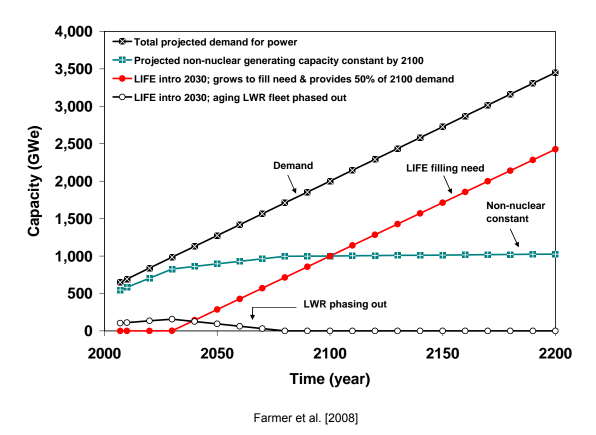


Figure 61 – Electrical generating capacity for 2nd scenario: LIFE introduced in 2030 to fill the projected need for electrical power.

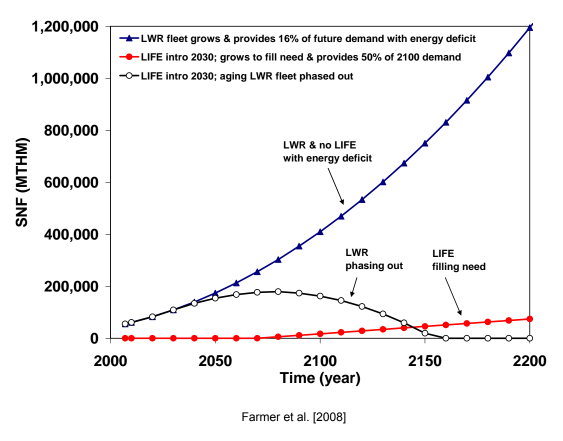


Figure 62 – SNF generated for two scenarios: (1) light water reactors with no LIFE engines and (2) introduction of LIFE engines in 2030.

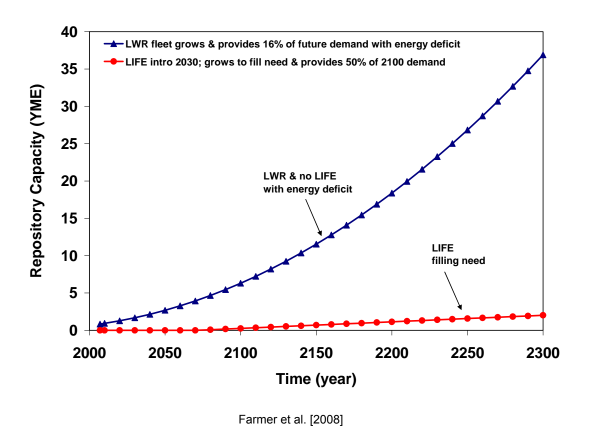


Figure 63 – Repositories required for two scenarios: (1) LWR with no LIFE and (2) LIFE intro in 2030.

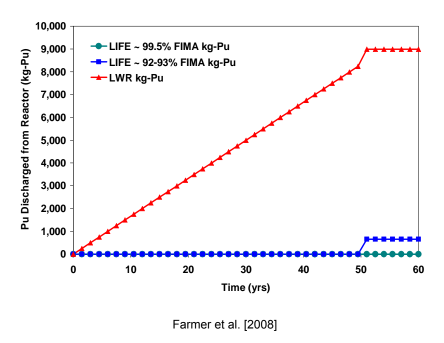


Figure 64 - LIFE engine's more complete burn generates far less plutonium per reactor than typical LWR.

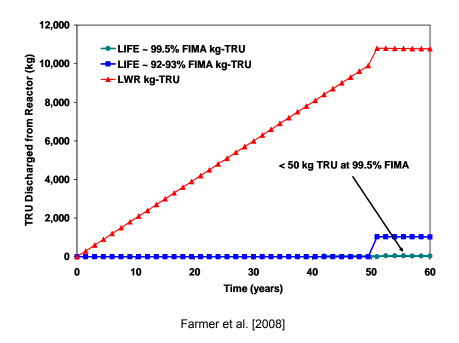
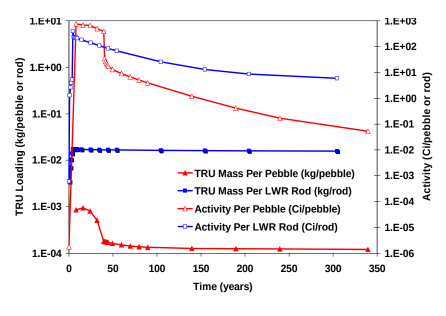
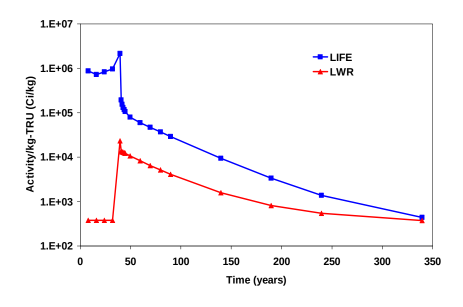


Figure 65 – LIFE engine's more complete burn also generates less TRU per reactor than LWR.



Farmer and Zhao [2008]

Figure 66 - LIFE fuel pebble is less attractive target than LWR fuel rod since the pebble has less TRU and more radioactivity.



Farmer and Zhao [2008]

Figure 67 – LIFE fuel (pebble) is a much less attractive target for theft than LWR fuel (rod): less TRU and more radioactivity.

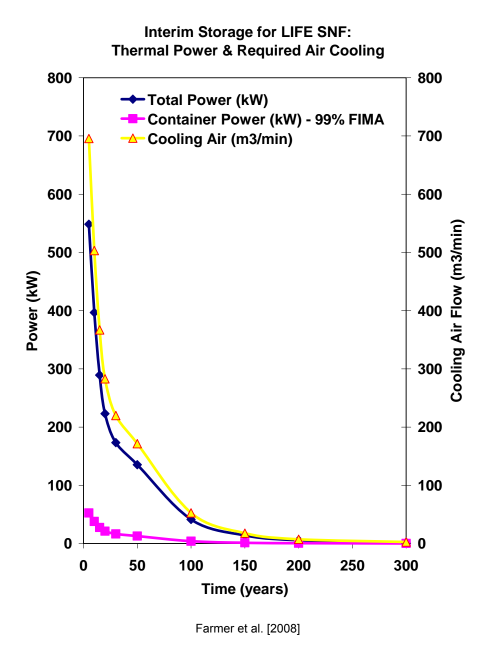


Figure 68 – Power Curves for interim storage of LIFE spent fuel in standard transportation, aging and disposal container during dry interim storage with area fraction for heat transfer between TRISO pebbles of 0.10.

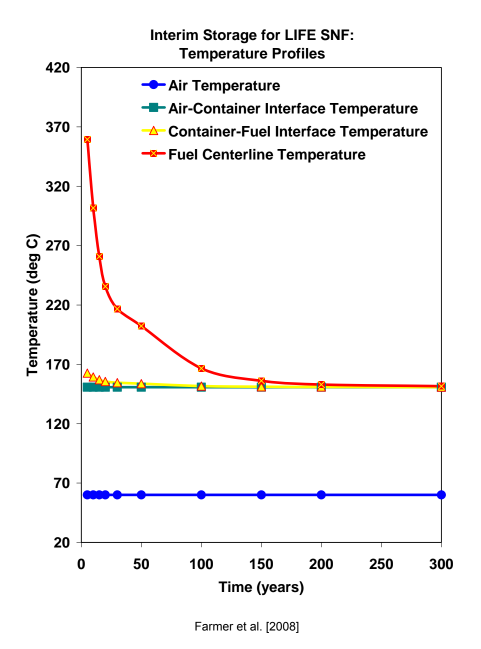
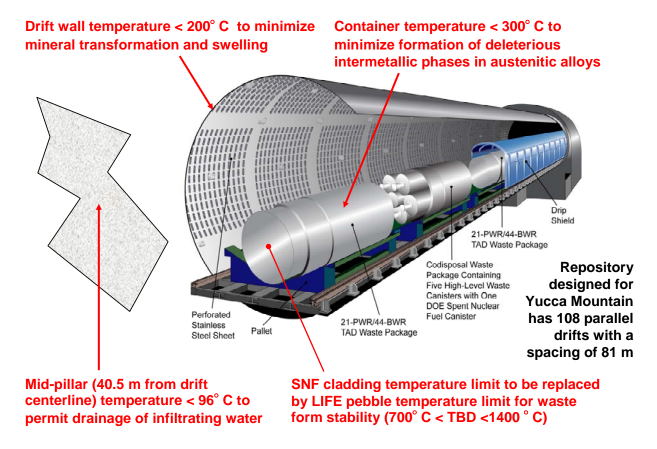


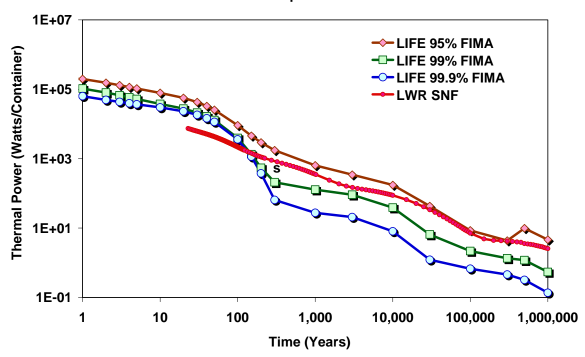
Figure 69 – Temperature and cooling air required for LIFE spent fuel in standard transportation, aging, and disposal container during dry interim storage with area fraction for heat transfer between TRISO pebbles of 0.10.



The repository design for Yucca Mountain has 108 parallel drifts with a centerline spacing of approximately 81 meters. Waste packages filled with spent fuel from LIFE satisfy repository temperature limits. For example, the drift wall temperature must be kept below 200°C to minimize deleterious mineral transformations and swelling. The midpillar temperature, at a distance of 40.5 meters from the drift centerline, must be kept below 96°C to permit drainage of the infiltrating water. The container, which will be fabricated from austenitic nickel-based Alloy C-22 must be maintained below 300°C to minimize formation of deleterious (P, σ and μ) intermetallic phases which deplete the matrix of those alloying constituents (Cr, Mo and W) which impart outstanding corrosion resistance. In the case of LWR SNF, the temperature limit for the zircaloy cladding is approximately 350°C. In the case of LIFE SNF, this would be replaced by the upper temperature limit for the TRISO fuel, which is assumed to lie between approximately 700 and 1400°C [U.S. DOE OCRWM 2008].

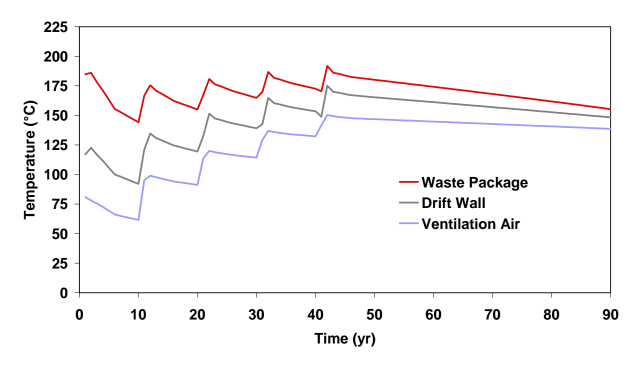
Figure 70 – Artist's rendering of drift inside of Yucca Mountain Repository with temperature limits shown.

Comparison of LIFE & LWR SNF Containers: Thermal Power per SNF Container



Thermal power of LIFE SNF decreases with increasing burn-up (%FIMA). The difference in FIMA power is burnup of actinides; fission products are in quasi-equilibrium. The higher LIFE thermal power can be accommodated by repository design options. Linear power density in the repository (kW/m) decreases with increasing burn-up (FIMA) [Blink et al. 2008].

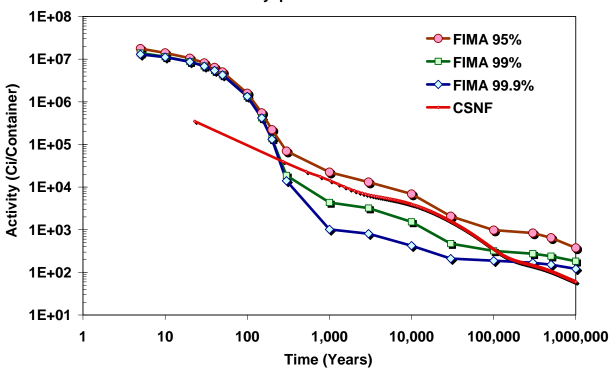
Figure 71 – A comparison of the heat generated by containers filled with LIFE and LWR spent nuclear fuels during the first million years.



Ventilation will be used to keep the temperature of LIFE SNF containers within limits during the pre-closure period of the repository, in a manner analogous to that for LWR SNF. This prediction of LIFE SNF container (waste package) temperature is for containers located near the drift's ventilation air exit. The ventilation air flow is assumed to be approximately 15 cubic meters per second (15 m³/s), with a velocity in the driftrs of approximately 1.79 miles per hour (1.7 mph). The saw-tooth pattern in the temperature-time profile is due to the phased, sequential periodic emplacement of the containers from the LIFE engine every 10 years. The burn-up is assumed to be approximately 99% FIMA [Blink et al. 2008].

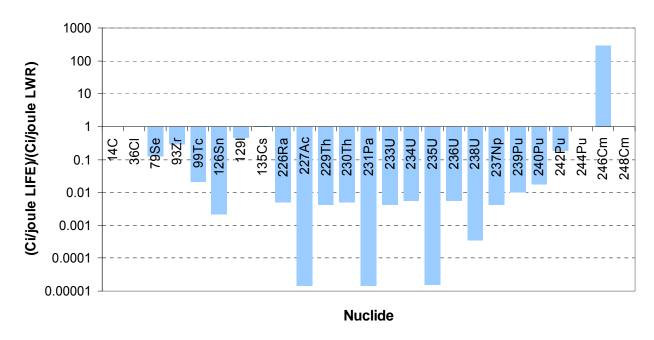
Figure 72 – Temperature of the container filled with LIFE spent fuel (waste package), drift wall, and ventilation air used for cooling during the pre-closure sequential emplacement of finge containers (from single LIFE engine) in a repository drift, assuming no preceding period of interim storage for cooling.

Comparison of LIFE & LWR SNF Containers: Activity per SNF Container



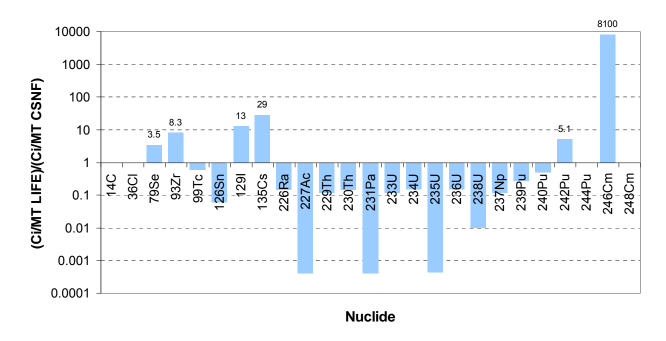
The radioactivity of LIFE SNF has also been modeled and decreases as the burn-up (%FIMA) increases. The difference in FIMA power is burnup of actinides; fission products are in quasi-equilibrium. Estimates for LIFE include some radionuclides (RNs) not included in the available estimates for the commercial spent nuclear fuel (CSNF). For example, Cm-246 is more important in LIFE SNF than in LWR SNF due to the high neutron fluence in LIFE (8 n captures). Other RNs such as Zr-93 and Nb-93m need to be closely scrutinized to determine why they are not included in available estimates for CSNF [Blink et al. 2008].

Figure 73 – A comparison of the radioactivity from containers filled with LIFE and LWR spent nuclear fuels during the first million years.



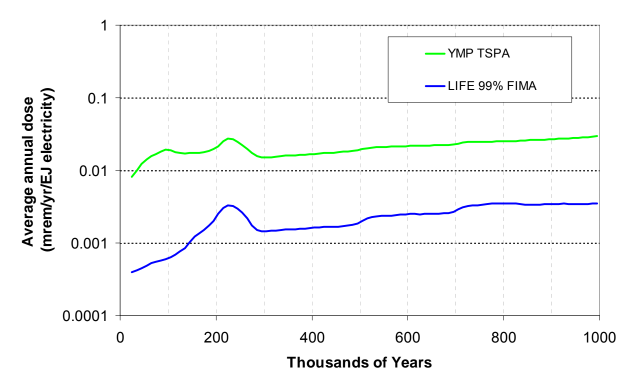
Plotted is the maximum ratio (at any time) of the activities per unit electrical energy generated by LIFE SNF (DU, 99% FIMA) and LWR SNF (41.2 GWt-day per metric ton $\sim 4.3\%$ FIMA) [Shaw et al. 2008].

Figure 74 – Comparison of energy-normalized inventories of radionuclides considered to be significant for the performance of the Yucca Mountain repository.



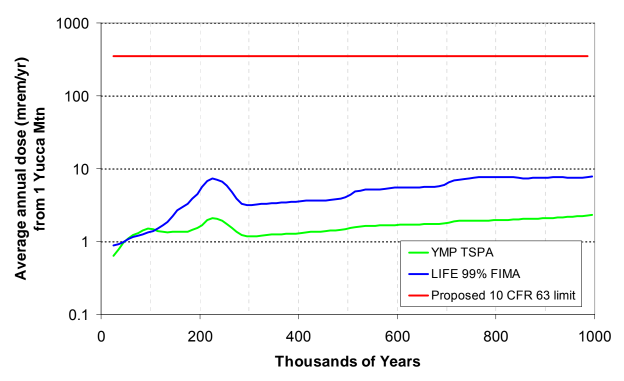
Plotted is the maximum ratio (at any time) of the specific activities (Ci/MTIHM) of LIFE SNF (DU, 99% FIMA) and LWR SNF (41.2 GWt-day per metric ton ~ 4.3% FIMA). Most significant nuclides considered significant in the Yucca Mountain TSPA are less abundant in LIFE SNF, but several are higher. Some nuclides in LIFE SNF are not accounted for in the Yucca Mountain TSPA (93 Zr, 244 Pu, 246 Cm, 248 Cm) [Shaw et al. 2008].

Figure 75 – Comparison of mass-normalized inventories of radionuclides considered to be significant for the performance of the Yucca Mountain repository.



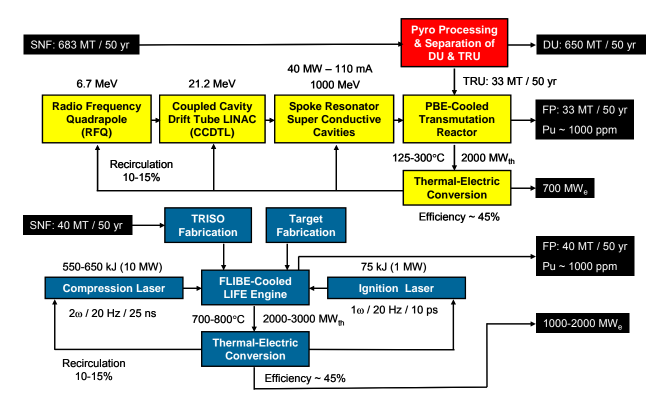
The dose predicted by performance assessment that is attributable to the generation of a given quantity of electricity is approximately ten times (\sim 10×) less for LIFE SNF than for LWR SNF [Shaw et al. 2008].

Figure 76 – The risk-to-benefit ratio for a Yucca Mountain repository completely filled with spent nuclear fuel from a fleet of LIFE engines compared to that of a similar repository completely filled with spent fuel from a fleet of light water reactors, showing risk for the LIFE scenario.



Scaled dose from a Yucca-Mountain-like repository containing LIFE SNF is of the same order of magnitude as the Yucca Mountain TSPA-LA results, and well below the proposed regulatory limit. Major contributors to long term dose from LIFE waste are ¹²⁹I, ¹³⁵Cs, and ²⁴²Pu. The LIFE Project will need to do performance assessment calculations using gualified and accepted software tools such as GoldSim in the future [Shaw et al. 2008].

Figure 77 – Simple scaling suggests that the risk of a Yucca Mountain repository completely filled with spent nuclear fuel from LIFE engines is slightly more than that of a repository completely filled with spent nuclear fuel from a fleet of light-water reactors, but still well within the statutory limits established with 10 CFR 63.



Another sub-critical system that has been proposed for the burning of spent nuclear fuel is known as accelerator transmutation of waste (ATW). In this process, reprocessing is used to separate TRU from LWR SNF for burning in a lead-bismuth cooled reactor. In contrast, SNF can be burned in LIFE without significant isotopic enrichment or chemical reprocessing [Farmer 2008].

Figure 78 - A side-by-side comparison of two proposed fuel cycles, one for the accelerator transmutation of waste and another with LIFE engines.

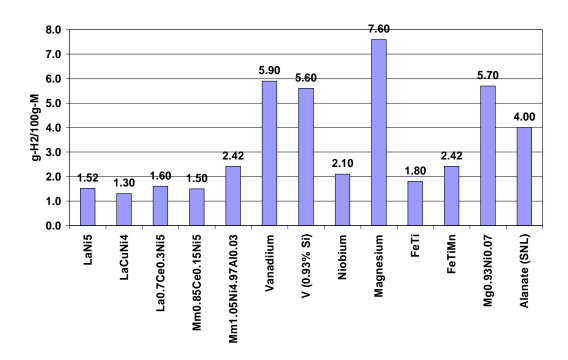
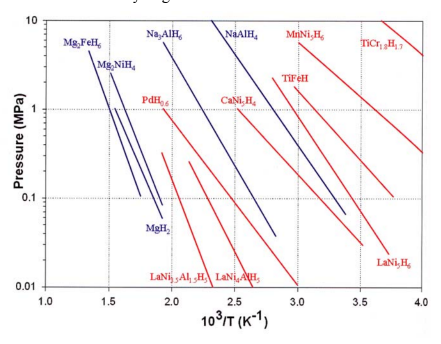


Figure 79 – LIFE requires safe on-site tritium storage with solid-state tritide systems, which have been demonstrated on relatively large scale.



Solid-state tritium storage media must be selected that will release the tritium with relatively mild temperature swings near ambient temperature. In Mg-Ni system may be an attractive system, with both high storage capacity, and modest operating temperature.

Figure 80 – Equilibrium pressure as a function of temperature for numerous solid-state storage media for hydrogen isotopes including tritium.

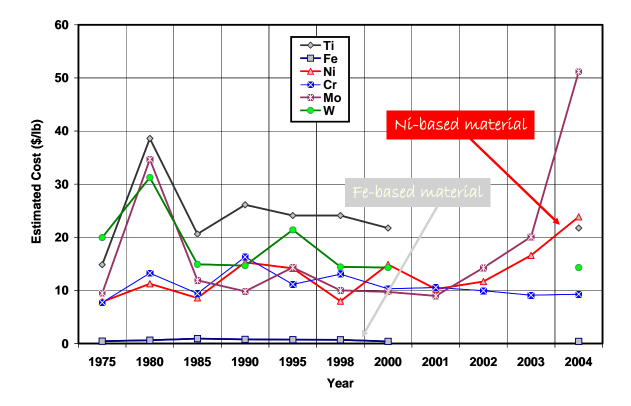


Figure 81 - Cost of raw materials required for the production of ferritic and austenitic steels, nickel-based alloys, and various refractory alloys. These costs were provided by the United States Geological Survey (USGS).

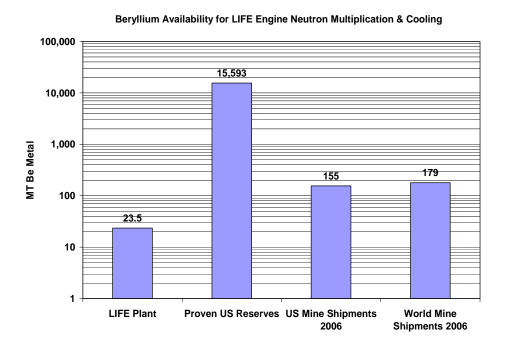


Figure 82 – The proven U.S. reserves, U.S. mine shipments, and world mine shipments of beryllium expressed in metric tons.

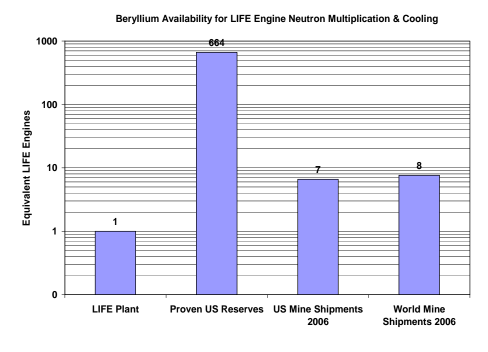


Figure 83 – The number of LIFE engines that can be built from proven US reserves, US mine shipments, and world mine shipments of beryllium.

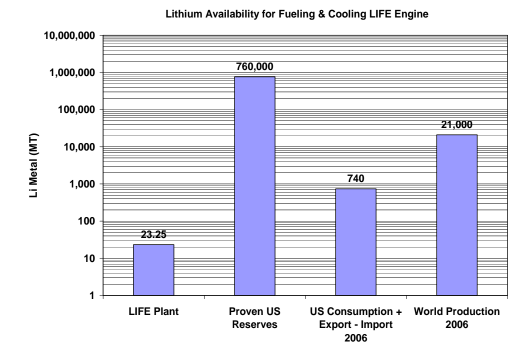


Figure 84 – The proven U.S. reserves, U.S. consumption and exports, deducting imports, and world production of lithium, expressed in metric tons.

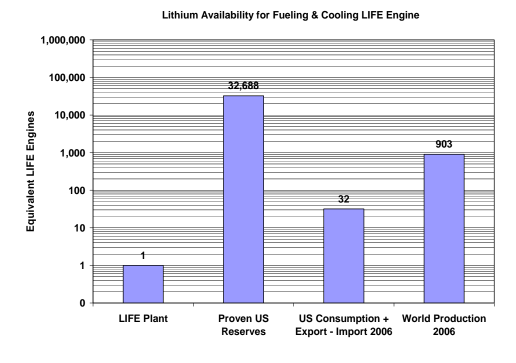


Figure 85 – The number of LIFE engines that can be built from proven U.S. reserves, U.S. consumption and exports, deducting imports, and world production of lithium.

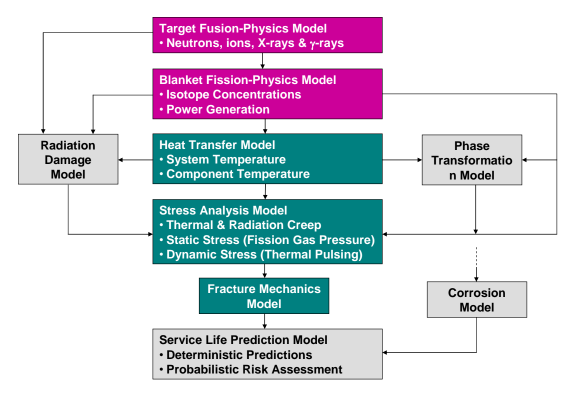


Figure 86 – A systematic approach to materials modeling will enable design of next-generation materials.

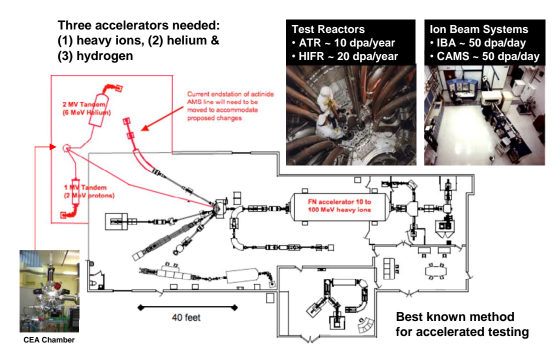


Figure 87 – Accelerated testing of materials will be done in three-beam accelerator.

Appendix 6 - Other High-Temperature Building Blocks

In the future, new high-performance first-wall composite materials could exploit the high melting points, and low vapor pressures of a broad range of known compounds. These compounds fall into several broad classes, including a wide variety of carbides, nitrides, oxides, intermetallics, silicides, as well as several refractory metals and alloys, some of which have already been discussed. Several refractory alloys have been explored for possible application as first-wall materials in fusion reactors [Ursu 1985; patent literature]. Several specific alloy compositions are given below.

Carbides

- 3500-4000°C: (Ta,Cr,Zr)C, HfC, Tac, ZrC, NbC
- 3000-3500°C: Ta₂C, TiC
- 2500-3000°C: SiC, VC, W₂C, MoC, ThC₂, WC
- 2000-2500°C: B₄C, Al₄C₃
- 1500-2000°C: Te₃C

Nitrides

- 3000-3500°C: HfN, TaN, BaN
- 2500-3000°C: ZrN, TiN, UN, ThN
- 2000-2500°C: AlN, Be3N, NbN, VN

Borides

- 3000-3500°C: HfB, TaB₂, ZrB₂, NbB₂
- 2500-3000°C: TiB₂, Ta₃B₄
- 2000-2500°C: VB₂, TaB, WB, W₂B₅, TiB, MoB, CrB₂, MoB₂, CrB, Ta₃B₂, TiB, Mo₂B

Intermetallics

- 3000-3500°C: Re₅W₂
- 2500-3000°C: MoW
- 2000-2500°C: CrAl, Mo₃Al, UBe₂, Zr₅Sn₃
- 1500-2000°C: Cr₃Ta, NiAl

Silicides

2500-3000°C: Ta₃Si

• 2000-2500°C: TaSi₂, W₅Si₃, Zr₂Si, WSi₂, ZrSi, V₃Si, Mo₃Si, Mo₄Si, MoSi₂

Refractory alloys are of particular interest for fusion and fusion-fission hybrid reactors due to: relatively high melting points, which enable this type of material to maintain high strength at elevated temperature. Refractory alloys are much more formable than ceramics. Refractory metals are found in Groups IVB, VB and VIB of the periodic table, and include Nb, Ta, Cr, Mo, and W. Specifically, these materials include:

Carbon

The favorable high-temperature properties of graphite have led to its prolific use in aerospace applications. The graphitic form consists of parallel planes of carbon atoms, arranged in hexagonal cells. This unique structure enables graphite to form intercalation compounds with lithium cations (Li+). The tendency of graphite to from intercalation compounds with the lithium cations in FLIBE and FLINABE would prove problematic in a fusion-fission hybrid reactor. This well-known characteristic has led to the use of graphite as the preferred cathode material in virtually all modern high-performance rechargeable Li-ion batteries.

• AW 12.011, AN 6, MP 3367°C, SUB 3367±25°C, BP 4827°C

• Amorphous : $\rho = 1.8 \text{ to } 2.1 \text{ g/cm}^3 (25^{\circ}\text{C})$

• Graphite: $\rho = 1.9 \text{ to } 2.3 \text{ g/cm}^3 (25^{\circ}\text{C})$

• Diamond: 3.15 to 3.5 g/cm³ (25°C)

Tungsten

Tungsten is also known as Wolfram. It is a true metal with the highest melting point and lowest vapor pressure; tantalum-carbide/graphite composite melts at 6760 °C. It forms oxides that are relatively insoluble in acid, and is one of the alloying elements added to austenitic nickel-chromium-molybdenum alloys to stabilize passive film in acidic oxidizing regions of the Pourbaix diagram, and significantly increases the pitting resistance equivalence number (PREN) of these alloys. In regard to the current nuclear fuel cycle, it is a key ingredient of nickel-based Alloy C-22 that will be used to construct 11,000 disposal containers for the Nation's 70,000 MTU spent nuclear fuel.

- AW = 183.85, AN 74, MP 3410 \pm 20°C, BP 5660°C, ρ = 19.3 g/cm³ (20°C)
- Ore: wolframite [(Fe,Mn)WO₄], scheelite [CaWO₄], huebnerite [MnWO₄], ferberite [FeWO₄]
- Sources: California, Colorado, South Korea, China, Bolivia, Ukraine/Russia

Tantalum

- AW = 180.95, AN 73, MP 2996°C, BP 5424 \pm 100°C, ρ = 16.654 g/cm³ (20°C)
- Ore: columbinte-tantalite [(Fe,Mn)(Mn,Ta)₂)₆]
- Sources: Zaire, Brazil, Mozambique, Thailand, Portugal, Nigeria, Canada

Molybdenum

This material is a byproduct of copper and tungsten mining; many ultra-high strength steels with yield strengths above 300,000 pounds per square inch (psi) contain this element at 0.25 to 8 weight percent. This alloying element is also added to austenitic stainless steels and nickel-chromium-molybdenum alloys (Type 316L Stainless, HastelloysTM and InconelsTM) to stabilize passive film in acidic oxidizing regions of the Pourbaix diagram, and significantly increases the pitting resistance equivalence number (PREN) of these materials. Even more important than tungsten in regard to its role in high-performance nickel-based alloys, it is a key ingredient of nickel-based Alloy C-22. In regard to the current nuclear fuel cycle, high-molybdenum Alloy C-22 will be used to construct 11,000 disposal containers for the Nation's 70,000 MTU spent nuclear fuel; sulfide serves as solid lubricant; catalyst; essential trace element for plant nutrition. Due to the entry of the Chinese into the world metals market, the costs of such high-performance alloys have skyrocketed. Recent procurements of Alloy C-22 were at a cost of \$37 per pound, driven significantly by the cost of both nickel and molybdenum.

- AW = 95.94, AN 42, MP 2617°C, BP 4612°C, $\rho = 10.22 \text{ g/cm}^3 (20^{\circ}\text{C})$
- Ore: molybdenite [MoS₂], wulfenite [(PbMoO₄)₃], powellite [Ca(Mo,W)O₄]
- Sources: relatively plentiful

Molybdenum Alloys

- 99.4%Mo-0.5%Ti-0.08%Zr-0.01%C
- 93%Mo-7at.%Re to 70%Mo-30at.%Re

Niobium

Niobium is also known as Columbium, and is unique in that it has superconductive properties.

- AW = 92.91, AN 41, MP 2468 \pm 10°C, BP 4742°C, ρ = 8.57 g/cm³ (20°C)
- Ore: associated with carbonatities; niobite-tantilite; pyrochlore; euxenite
- Sources: Canada, Brazil, Nigeria, Zaire, Ukraine/Russia

Niobium Alloys

• 81%Nb-10%Mo-3%Ti-1%Zr

Vanadium

Vanadium forms refractory alloys that have been shown to have reasonably good corrosion properties in the presence of liquid lithium, FLIBE and FLINABE. This element is plentiful, and is found in over 65 different minerals. Its ores are frequently associated with phosphate rock and iron ore.

- AW = 50.94, AN 23, MP 1890 \pm 10°C, BP 3380°C, ρ = 6.11 g/cm³ (18.7°C)
- Ore: carnotite, roscoelite, vanadinite, patronite
- Sources: relatively plentiful

Vanadium Alloys

- 80%V-20%Ti
- (81-94)%V-(3-9)%Cr-(3-10)%Ti
- V-4Cr-4Ti

Chromium

Chromium is produced through the chemical reduction of oxide ore with metallic aluminum, which is produced electrolytically. From an environmental point of view, chromium is paradoxical: hexavalent chromium is carcinogenic, while trivalent chromium is essential for human life. Chromium is the element primarily responsible for passive film formation on iron-based 300-series stainless steels, including Types 304L and 304L. The passive film formed from chromium is stable over modest ranges of potential and pH. Due to its role in the passivity of steels, it is also a key ingredient of nickel-based Alloy C-22. As previously discussed in regard to tungsten and molybdenum, Alloy C-22 will be used to construct 11,000 disposal containers for the Nation's 70,000 MTU spent nuclear fuel.

- AW = 51.996, AN 24, MP 1857 \pm 20°C, BP 2672°C, ρ = 7.18 g/cm³ (20°C)
- Ore: chromite [FeCr₂O₄]
- Sources: Southern Rhodesia, Russia/Ukraine, Turkey, Iran, Albania, Finland, Madagascar, Philippines

Platinum

Platinum occurs in native form as silvery-white metal, and usually found with lesser amounts of iridium, osmium, palladium, ruthenium, and rhodium. This white precious metal recognized by pre-Colombian men in the Americas. Platinum is unique in its outstanding catalytic and electrocatalytic properties, and is the catalyst of choice in proton exchange membrane fuel cells for both oxygen cathodes and hydrogen anodes, at loadings of 0.5 to 2.0 mg/cm². As the world moves towards a hydrogen economy, the demands and costs are likely to rise significantly, until alternative electrocatalysts are created through bandgap engineering.

- AW = 195.09, AN 78, MP 1772°C, BP 3827 \pm 100°C, ρ = 21.45 g/cm³ (20°C)
- Ore: metal [native form]; sperrylite [PtAs₂]
- Sources: Colombia, Canada, Western United States, Others

Titanium

The National Strategic Reserve of Titanium Metal is located in Henderson, Nevada, at TIMET, Incorporated. At the present time, the Department of Defense is investing heavily in processes to achieve dramatic reductions in the cost of titanium, by shifting from chemical to electrolytic reduction, as was done for aluminum production in the early 20th century. It is frequently alloyed with small amounts of palladium to enhance its environmental resistance. Approximately 12.1 weight percent of Moon Rocks collected during Apollo 17 mission consisted of rutile (TiO₂), and most white paint pigment consists of rutile, which was used as a safe alternative to lead oxide. This particular oxide has unique photocatalytic properties. Titanium is strong lightweight metal used for many aerospace and defense applications. It is extremely corrosion resistant in saline environments, and is one of the materials of choice in desalination plants. In regard to the current nuclear fuel cycle, more than 10,000 titanium alloy (Ti Grade 7) drip shields are planned for emplacement in Yucca Mountain to protect waste packages from rock drop and dripping ground water, and will cost several billion dollars.

- AW = 47.90, AN 22, MP $1660\pm10^{\circ}$ C, BP 3287° C, $\rho = 4.54 \text{ g/cm}^3 (20^{\circ}\text{C})$
- Ore: rutile, ilmenite, sphene
- Sources: relatively plentiful

ⁱ. J. R. Engel, W. R. Grimes, W. A. Rhoades, J. F. Dearing, "Molten salt reactors for efficient nuclear fuel utilization without plutonium separation," ORNL/TM-6413 (1978).